Magnetization arrangement of hard magnetic phases and mechanism of magnetization and reversal magnetization of nano-composite magnets

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Abstract: During the process of directional solidification, laser remelting/solidification in the layer on sintered magnets, die-upsetting of cast magnets, or die-upsetting of nano-composites, the arrangements of the easy-magnetization-axes of the hard magnetic phases (Nd_2Fe_14B, SmCo_5 or Sm_2Co_17 type) in their designed directions have been studied. In Fe-Pt nano-composite magnets, attempts have been taken to promote phase transformation from disordered, soft magnetic $A^1$ to ordered, hard magnetic $L_{10}$ FePt phase at reduced temperatures. The dependence of the magnetization and reversal magnetization processes on the microstructures, involving the morphology and three critical sizes of particles of the FePt nano-composite magnets, are summarized. With the decrease of the nominal thickness of the anisotropic FePt film epitaxially grown on the single crystal MgO (001) substrate, the reversal magnetization process firstly changes from full domain wall displacement to partial magnetic wall pinning related to the morphology change, where the coercive force increases abruptly. The reversal magnetization process secondly changes from magnetic wall pinning to incoherent magnetization rotation associated with the particles being below the first critical size at which multi-domain particles turn into single domain ones, where the coercive force is still increased. And the reversal magnetization mode thirdly changes from incoherent to coherent rotation referred to the second critical size, where the increase of the coercive force keeps on. However, when the particle size decreases to approach the third critical size where the particles turn into the superparamagnetic state, the coercive force begins to decrease due to the interplay of the size effect and the incomplete ordering induced by the size effect. Meanwhile, due to the size effect, Curie temperature of the ultra-small FePt particles reduces.

Key words: permanent magnets; arrangement of easy-magnetization-axis; hard magnetic phases; nano-composite magnets; mechanism of magnetization and reversal magnetization; $L_{10}$ FePt; ordering; coercivity; size effect

1 Introduction

Until 2003, researches on nano-composite magnets have made little progress, such as their energy products for more than a decade still being 159–199 kJ/m³, lower than one fifth of the theoretic value of about 1 MJ/m³ of Nd-Fe-B nano-composite magnets[1–2]. One of the reasons lies in the fact that the magnetization arrangement of the hard magnetic phases has not been aligned. The studies in Refs.[2–5] indicate that the arrangement of the easy magnetization axis [001] of RE_2Fe_14B is the key issue in improving the magnetic properties of the nano-composite magnets. With the advancement in the arrangement, their energy product has reached 437.8 kJ/m³[3]. But their coercive forces are relatively low due to the decay of coercivity during hot-working process[2–6]. The mechanisms of the magnetization process and reversal magnetization process of the nano-composite magnets are distinctly different from those of the magnets with grains in micron size[7]. Thus, it is essential to explore the effects of microstructures, mainly the morphology and particle sizes, on the magnetization and reversal magnetization process which should directly associate with the coercive force of nano-composite magnets.

In this work, two key topics are mainly reviewed. One is the arrangement of the easy magnetization axes of hard magnetic phases during new synthesis routs, although some of them are in micron size. And the other...
is the mechanism of the magnetization and reversal magnetization of nano-composite magnets by the use of FePt based thin films or nano-particles since their microstructures are more easily controlled than bulk nano-composite magnets, in the expectation of improving the magnetic properties of nano-composites, such as Nd$_2$Fe$_{14}$B/α-Fe, Nd$_2$Fe$_{14}$B/Fe$_3$B, Sm$_2$Co$_{17}$/Co, and Sm$_1$Co$_5$/Co.

2 Easy-magnetization-axis arrangement of Sm$_2$Co$_{17}$, Sm$_1$Co$_5$ and RE$_2$Fe$_{14}$B

In order to fully make use of the high saturation magnetizations of rare earth transition metal compounds, such as Sm$_1$Co$_5$, Sm$_2$Co$_{17}$ and Nd$_2$Fe$_{14}$B, and to obtain high magnetic properties, it is necessary to prepare bulk magnets with the arrangement of the easy magnetization axes of the hard magnetic phases in the whole samples in one designed direction during a certain new synthesis process, such as directional solidification, laser remelting/solidification, cast/die-upset, and hot-pressing/die-upset.

2.1 Directional solidification of Sm-Co alloys

In the Sm-La-Co-Cu-Fe-Zr system with potentially high magnetic properties, the microstructures of directionally solidified rods are shown in Fig.1. The matrix is (Sm,La)$_2$(Co,Fe,Cu,Zr)$_{17}$; the other magnetic phase is (Sm,La)$_1$(Co,Cu,Fe)$_5$; and the dendrites are composed of cobalt phase[8–11]. When the crystal growth velocity is decreased from 21 to 7 mm/min, or the ratio $G/R$ ($G$ is the temperature gradient at the solid/liquid interface, and $R$ is the crystal growth rate) is increased, the arrangements of the morphology of these three kinds of phases are changed from about 45˚ (Fig.1(a)) to 0˚ (Fig.1(c)) (the angle between the solidification direction and the axes of the dendrites). Although the columns of the hard magnetic phases and the cobalt dendrites are all parallel to the solidification direction (seen in Figs.1(b) and (c)), the magnetic domain

![Typical microstructures of Sm-La-Co-Cu-Fe-Zr system directionally solidified at different crystal growth rates on longitudinal section[11]: (a) 21 mm/min; (b) 15 mm/min; (c) 7 mm/min; (d) Maze-like domains](image-url)
walls are not always consistent with the solidification morphology (Fig.1(b)). The typical magnetic domain walls on the magnetic phases are of about 20° with the constrained solidification direction, or the rod axis. The magnetization arrangement angle, $\alpha$, is defined as the angle between the constrained solidification direction and the macroscopic magnetization axis of whole sample. As seen in Fig.1(d), there are maze-like domains on the longitudinal section, and $\alpha$ is equal to 90°. The alloy compositions of the sample in Fig.1(d) are different from those in Figs.1(a), 1(b) and 1(c).

If the angle $\alpha$ of a sample is about 20°−90°, the statistics results show that the remanence is 0.09−0.37 T, which is too low to have any practical use. But the samples with $\alpha$ of 0°−5°, directly synthesized by the directional solidification (seen in Fig.2), have very high remanence, 0.85−1.20 T, the same as the anisotropic counterpart prepared directly by the traditionally sintered method, with particles magnetized in a field above 800 kA/m.

In Sm-La-Co-Cu-Fe-Zr system, it is discovered that, whether the angle equals 0°−5° or does not depends not only on compositions but also on the $G/R$ ratio under certain conditions. For some alloys, such as Co-32.0Sm-3.5La-15Fe-6.5Cu-2.5Zr (mass fraction, %), if the gradient $G$ at solid/liquid interface is 200 K/cm, the angle can be changed from 90° to 0° as the growth rate is decreased from 12 mm/min to 6 mm/min. The reason maybe lies in the fact that the ratio $G/R$ is higher than the critical value, above which not only the morphologies of the magnetic phases but also the magnetic domains are parallel to the constrained solidification direction.

For other alloys, even both the temperature gradient and crystal growth rate are changed, $\alpha$ is not 0°−5° under the conditions of directional solidification process of 1 553−1 723 K and crystal growth rate of 0.3−25 mm/min. Only by sintering method, can their micron particles be arranged in a magnetic field and the remanences of the sintered samples be high enough as magnets. It is deduced that the atoms in these magnets occupy error points in the crystal lattice of (Sm,La)$_3$(Co,Cu,Fe,Zr)$_7$ or (Sm,La)$_3$(Co,Cu,Fe)$_5$, during the non-equilibrium directional solidification process. So, these alloys cannot be now prepared directly by the solidification process[10−11].

For the alloys with the angle equal to 0°−5°, the effects of heat treatment on magnetic properties have been studied, and these magnets, finally with 143.3 kJ/m$^3$, directly prepared by directional solidification and heat treatment, are more ductile than the sintered ones with the same compositions.

However, the macroscopic $c$-axis texture orientation of Nd$_2$Fe$_{14}$B cannot be formed directly through directional solidification under the conditions of temperature gradient at the solid/liquid surface of 200−1 000 K/cm, and the crystal growth rate of 12−250 μm/s[12−13].

2.2 $c$-axis texture of Nd$_2$Fe$_{14}$B solidified in laser remelting/solidification layer on anisotropic magnets

Three kinds of commercial anisotropic sintered Nd$_{15}$Fe$_{77}$B$_8$ magnets with their magnetization direction along $X$-, $Y$-, and $Z$-axis, respectively, have been cut into cube, whose dimensions are 42 mm×20 mm×8 mm. Their surfaces parallel to XOY plane, are remelted/solidified by 5 kW Roffin-Sinar 850 type of CO$_2$ laser, whose beam direction is in the opposite $Z$-axis. The above process is under argon protection. The scanning parameters, such as the power 2.5 kW, the diameter of
the laser light spot 0.05 mm, and the scanning velocity 10–30 mm/s are used.

Only when the magnetization axis of the whole anisotropic magnet substrates is perpendicular to the heat flux direction of the layer, can c-axis texture be obtained in the laser melting/solidification layer. The same orientation of the c-axis texture of Nd$_2$Fe$_{14}$B in the layer can be achieved as that of the anisotropic substrates (seen in Fig.3). The plate-like magnet domain walls in the layer are parallel to those in the substrates. If the geometric relationship between the magnetization axis of the substrates and the heat flux direction of the laser melting/solidification pool is different from the above, c-axis texture cannot be formed and the laser melting/solidification layer will be isotropic[14–18].

![Fig.3 Magnetic domains in longitudinal section in laser melting/solidification layer on anisotropic magnet with its magnetization axis normal to heat flux direction[18]](image)

### 2.3 [001] texture of cast/die-upset Pr$_{x}$Fe$_{93.5}$-$x$B$_{5}$Cu$_{1.5}$

Compared with the magnets prepared by powder sintering, the cast/die-upset process of RE-Fe-B-Cu magnets includes no powder preparing step. Thus, cast/die-upset magnets should exhibit lower oxygen content than the sintered ones[19]. However, the residual magnetization of 0.78–1.12 T, in the previous work, is much lower than that of sintered ones. Their highest intensity ratio of $I_{006}/I_{105}$ determined by X-ray diffraction is merely 0.76, indicating that the easy magnetization axes of Pr$_{x}$Fe$_{93.5}$B$_{5}$Cu$_{1.5}$ grains are poorly aligned[20].

The cast slabs are first cut into small pieces for die-upset. Die-upset is carried out at 973–1 173 K with strain rate of 10$^{-3}$–10$^{-1}$ s$^{-1}$, and strain of 0–80%[21].

When die-upset parameters are under the conditions of temperature 973–1 173 K, strain rate 10$^{-3}$ s$^{-1}$ and strain 80%, the magnetic domain walls (seen in Fig.4) on the longitudinal plane are parallel to the die-upset direction. And the intensity of $I_{006}$ of the die-upset Pr$_{x}$Fe$_{93.5}$-$x$B$_{5}$Cu$_{1.5}$ is the highest on the diffraction pattern. The residual magnetization of these magnets has reached 1.3 T[21]. It is confirmed that die-upset is in favor of obtaining the anisotropic RE$_2$Fe$_{14}$B bulk magnets.

![Fig.4 Magnetic domains of cast Pr$_{x}$Fe$_{93.5}$-$x$B$_{5}$Cu$_{1.5}$ die-upset at 973–1 173K, strain rate of 10$^{-3}$ s$^{-1}$ and strain of 80%, on longitudinal section[21]](image)

### 3 Hot-pressed and die-upset Nd-Fe-B nano-composite bulk magnets

#### 3.1 Arrangement of Nd$_2$Fe$_{14}$B phase along [001]

It is necessary to note that the die-upset process in the hot-pressing/die-upset method is different from that in the above cast/die-upset method. Only when its heating rate is much larger than that of the latter, and its die-upset duration at high temperature is much shorter than that of the latter, can the [001] texture be achieved in Nd-Fe-B nano-composite bulk magnets, and can their energy products much higher than 159.2–199 kJ/m$^3$, be reached.

In Fig.5[5], it is observed from the XRD diffraction pattern, hot-pressed Nd$_{12}$Dy$_2$Fe$_{73.2}$Co$_6$Ga$_{0.6}$B$_{5.6}$ alloy consists of randomly oriented Nd$_2$Fe$_{14}$B grains with homogeneous distribution in the grain size of 50–100 nm, whose (006) and (008) peaks are inconspicuous while (314), (410) and (214) appear as the main reflection planes, somewhat similar to the XRD pattern of the same composition powders, indicating that the hot-pressed magnet has isotropic property. While the microstructures of the die-upset alloy(seen in Fig.5(b)) show that the Nd$_2$Fe$_{14}$B grains in the shape of platelet with 50–100 nm in thickness and 200–500 nm in length are well aligned perpendicular to the press direction. Its XRD pattern shows that (006) peak becomes the dominant reflection, (004) and (008) peaks are obvious(shown in Fig.5(a)), indicating the formation of magnetic texture by the alignment of the easy-magnetization-axis of tetragonal unit cell of the Nd$_2$Fe$_{14}$B phase after the hot-pressing/die-upset process.

#### 3.2 Influence of die-upset parameters on magnetic properties

As seen in Fig.6[4], with the increase of deformation strain, the values of $B_r$ and $(BH)_{\text{max}}$ of the
Nd$_{13.6}$Fe$_{73.7}$Co$_{6.6}$Ga$_{0.6}$B$_{5.5}$ alloy increase monotonously, but $H_c$ decreases. Moreover, when the deformation strain is in excess of 65%, the magnetic properties will decay considerably, resulting from the crack present in the magnets[4].

$B_r$, $H_c$ and $(BH)_{max}$ reach their maximum values simultaneously after die-upset at 1 023 K in addition to the best squareness of loops, indicating that 1 023 K is the optimal temperature for the hot-working of Nd$_{13.6}$Fe$_{73.7}$Co$_{6.6}$Ga$_{0.6}$B$_{5.5}$ alloy. While die-upset temperature 1 273 K is so high that it will lead to excessive grain growing and coalescing, which results in the decay of $B_r$ and substantial decline of $H_c$[4].

$B_r$ is 1.2 T for the magnet die-upset of 873 K where the Nd-rich grain boundary is not in liquid state, indicating that the magnet with certain anisotropy can be obtained even without the grain boundary in liquid state during die-upset process. This phenomenon is worthy of studying further.

### 3.3 Effects of Nd content on magnetic properties

Fig.7 shows the effects of Nd content in Nd$_{x}$Fe$_{94-x}$B$_6$ on the magnetic properties of powder, hot-pressed magnet and die-upset one[6]. With the Nd content decreasing, a relatively high $B_t$ of the Nd$_{x}$Fe$_{94-x}$B$_6$ die-upset magnet is obtained to be about 1.2 T, for 12%−18% (molar fraction) Nd; then it steeply decreases for 10%−12% Nd; however, it increases to 0.8−1.0 T for 6%−10% Nd. From these results, $B_t$ is lower than 1.0 T for $x$(Nd)<11%. Although this Nd-lean alloy consists of Nd$_2$Fe$_{14}$B and $\alpha$-Fe phase, it is difficult to induce texture in this kind of magnet[6].

$H_c$ of the die-upset Nd$_{x}$Fe$_{94-x}$B$_6$ magnet decreases monotonously with the decrease of the Nd content. The
Fig. 7 Effects of Nd content on magnetic properties of powder (a), hot-pressed (b) and die-upset magnet (c) [6]

For Nd content, the magnetic properties decrease is slow for 14%-18% and 6%-12% Nd, but substantial for 12%-14% Nd (molar fraction). 

\[(BH)_{\text{max}}\] of the die-upset Nd$_2$Fe$_{14}$B magnet varies slightly for \(x(\text{Nd}) \geq 14\%\), and keeps a value of about 250 kJ/m$^3$ with the decrease of Nd content. It decreases drastically for 12%-14% Nd, and with further decrease of Nd content, \((BH)_{\text{max}}\) keeps constant.

Using the hybrid method (blending the powders of RE-rich Nd-Fe-B alloy and RE-lean one, even of both RE-rich alloy and soft-magnets without any RE content, then hot-pressing and die-upset), energy product of Nd$_2$Fe$_{14}$B based nano-composite magnets have reached 437.8 kJ/m$^3$[3].

With discussion above, anisotropy in the Nd$_2$Fe$_{14}$B nano-composite magnets could be induced by both the hot-pressing and die-upset process, accompanying with the increase of remanence and energy product but the decay of coercivity, which prevent the further improvement of energy product after annealing. The reversal magnetization and coercivity mechanism of bulk nano-composite magnets certainly depend on the microstructure details, such as the three-dimensional spatial distribution of the phases, and grain size. Thus, it is essential to establish the relationship between the microstructure and coercivity clearly. However, up to now, the coercivity mechanism of nano-composite magnets, Nd$_2$Fe$_{14}$B/$\alpha$-Fe or Nd$_2$Fe$_{14}$B/Fe$_3$B, has not been developed. So in this work, the mechanism of FePt-based nano-composite magnets is reviewed.

4 Ordering transformation and coercivity mechanism of FePt-based nano-composite magnets

The FePt thin films with easily controllable and reproducible microstructures are comfortable to be applied in the field of coercivity study. Especially, thin-film growth allows the control of layer thickness, and it can also provide a means for crystallographic alignment of the easy magnetization axis of the hard phase. An additional advantage of the FePt thin films is to allow a realistic estimation of the ultimate gain in performance that may potentially be realized in exchange-coupling bulk magnets. The coercivity study of FePt based nano-composite magnets is expected to guide the efforts on the enhancement of the magnetic properties of other kinds of nano-composite magnets with much lower material cost, such as Nd$_2$Fe$_{14}$B/$\alpha$-Fe, Nd$_2$Fe$_{14}$B/Fe$_3$B, Sm$_2$Co$_{17}$/Co, and Sm$_1$Co$_5$/Co.

$L1_0$ FePt alloy has attracted much attention for decades due to its remarkable uniaxial magnetocrystalline anisotropy ($K_u \approx 7 \times 10^6$ J/m$^3$) along the c-axis direction[22-23], which leads to good intrinsic coercive forces with $x=0.50-0.68$ in Fe$_x$Pt$_{1-x}$ (seen in Fig.8)[24], chemical stability, high Curie temperature ($T_c > 723$ K)[24], and attractive saturation magnetization ($4\pi M_s = 1.38$ T[23]) in dense bulk form. In recent years, thin films or particles of $L1_0$ FePt are expected to be applied in the synthesis of ultrahigh density magnetic recording media and micro-electro-mechanical systems (MEMS). According to KNELLER and HAWIG[25], hard phase and soft phase will be well coupled with each other when the size of the soft phase is within the critical one $b_{cs} = \frac{\pi n_0 J}{2K_u} = 10$ nm for $L1_0$ FePt, where $J$ is exchange stiff constant, resulting in a very high energy
In this work, the ordering transformation is firstly discussed for there are both soft and hard magnetic phases after ordering transformation.

4.1 Promoting ordering transformation at reduced temperature

The equi-atomic FePt compounds have two kinds of crystal structural phases: one is $A_1$ or fcc, which is chemically disordered and exhibits soft magnetic behavior; and the other is $L_{10}$ or fct, which is chemically ordered and magnetically hard. Since room-temperature as-synthesized FePt materials take the $A_1$ structure, heat-treatment (usually above 773 K) is required to convert the disordered $A_1$ to the ordered $L_{10}$. Excessive grain growth that is deleterious to coercivity easily takes place at elevated temperature, thus, it is desirable to find a way to produce $L_{10}$ FePt phase at reduced temperatures.

4.1.1 Ordering kinetics

The $A_1 \rightarrow L_{10}$ transformation is induced by the nucleation and growth of the ordered phase in the disordered matrix; and both are thermally activated processes[27]. The activation energy for nucleation($E_a$) and growth($E_g$) could be expressed respectively as

$$E_a = -k_B T \ln \frac{R_n}{R_{n0}}$$  \hspace{1cm} (1)

$$E_g = k_B T \ln \frac{t_E}{t_{E0}}$$  \hspace{1cm} (2)

where $k_B$ is the Boltzman’s constant; $T$ is the annealing temperature; $R_n$ is the nucleation rate; $t_E$ is the saturation time constant; $R_{n0}$ and $t_{E0}$ are constants; $E_a = [(0.5 - 0.6) \pm 0.1] eV$ and $E_g = (0.9 \pm 0.1) eV$ are estimated, respectively, according to experimental results[28]. $E_a$ is lower than $E_g$, resulting in an easy nucleation process for the $L_{10}$ ordered domains in the FePt thin films, and a difficult growth process at high ordering temperature. The activation energy for growth can be compared to that for atomic diffusion in disordered FePt matrix, thus, it is suggested that the growth of the $L_{10}$ ordered domains is determined by atom diffusion[28].

Based on this understanding, the addition of the third element, introduction of defects in films or between the films and buffer layers, using off-stoichiometric method, transformation under high pressure, processing in external magnetic field and with different kinds of sputtering gas could be effective to be applied to lowering down the ordering temperature.

4.1.2 Adding the third element

It could be found in Fig.9 that the ordering beginning temperatures where coercivity begins to steeply increase, resulting from the formation of $L_{10}$ phase with large $K_u$, are 598, 623 and 673 K for chemically-synthesized (FePt)$_{85}$Au$_{15}$, (FePt)$_{88}$Ag$_{12}$, and Fe$_{48}$Pt$_{52}$ nano-particles, respectively[29]. And the ordering temperatures are 823, 873 and 873 K, respectively, for (Fe$_{0.588}$Pt$_{0.412}$)$_{85}$Pd$_{15}$, (FePt)$_{90}$Cr$_{10}$ and (Fe$_{0.412}$Pt$_{0.588}$)$_{85}$Cu$_{15}$ nano-particles. The increase of the coercivity of Fe$_{48}$Pt$_{52}$ particles is the slowest, whose slope of $H_c-T$ curve is parallel to that of (Fe$_{0.412}$Pt$_{0.588}$)$_{85}$Cu$_{15}$. Compared with Fe$_{48}$Pt$_{52}$, additive Au and Ag decrease the ordering temperature by 75 K and 50 K, respectively; while additive Pd, Cr and Cu increase the ordering temperature by 150 K, 200 K and 200 K. For (FePt)$_{85}$B$_{15}$ film in thickness of 10 nm, the coercivity increases drastically above 673 K, while it takes place at about 873 K for the pure FePt film[23].

Decrease of the activation energy, i.e. ordering temperature, for Ag and Au-doped Fe-Pt nano-particles is attributed to the segregation of Ag and Au atoms from $A1$ FePt to form Ag and Au phases during the heat treatment, leaving the increased number of vacancies, then resulting in the increase of the diffusion coefficient of Fe and Pt atoms[30]. Somewhat similarly, decreasing
of the ordering beginning temperature by boron addition is attributed to the high diffusivities of Fe and Pt due to the defects created by the movement of the boron atoms to interstitial or substitutional lattice sites[29]. Thus, it can be concluded that reduced ordering temperature could be obtained by adding the third element which yields the defects to be beneficial to the diffusivities of Fe and Pt atoms.

4.1.3 Defects

It is well agreed that defects play an important role in the $A1\rightarrow L1_0$ transformation. Defects or internal stress may contribute to the formation of the crystalline nucleus, the grain boundaries and dislocations acting as diffusion tunnels[32]. And the addition energy, stored in the form of a variety of defects including grain boundaries, dislocations, internal strain or stress, will be released during annealing process, which is found to shorten the ordering transformation time and/or lower the ordering temperature of $L1_0$ alloys[33]. Fig.10 shows that the value of $S_0$, namely, the volume fraction of $L1_0$ CoPt phase which is used to describe the degree of ordering, is retarded by 0.1−0.25 for the preannealed films compared with the un-treated ones. This is interpreted to result from the phenomenon that the preannealing treatment eliminates mostly dislocations or vacancies.

4.1.4 Off-stoichiometric alloys

As shown in Fig.11, the size of $L1_0$ crystal cell, $a$, holds a constant for $x \leq 38$ in Fe$_x$Pt$_{100-x}$ films, then decreases monotonically when $x > 38$. On the other hand, the value of $c$ decreases monotonically when $x \leq 38$, considering that Fe sites in the $L1_0$ structure are partly replaced by Pt atoms[35]. It is worth noting that the degree of atomic ordering of the $L1_0$ FePt phase, quantitatively described by $S=0.85[|I_{001}/I_{002}|]^{1/2}$[36], is benefited from the decrease of $c/a$ and reaches a maximum (0.6±0.1) around $x=38$ in Fe$_x$Pt$_{100-x}$ films grown at just 573 K where the value of $c/a$ reaches the
minimum (0.955). It is expected that the effect of strain, resulting from the tetragonal c/a, is dominant in this course although the mechanism is not clearly understood now.

4.1.5 Lattice mismatch $\Delta a/a$ between Fe$_x$Pt$_{1-x}$ films and buffer layers

In Fig.12, it is found that $S$ increases monotonically for Fe$_{38}$Pt$_{62}$ films with $\Delta a/a$ increasing, and reaches the maximum value of 0.8±0.1 at $\Delta a/a=9.0\%$. On the other hand, for Fe$_{52}$Pt$_{48}$ films, when $\Delta a/a<5.4$, $S$ increases drastically and achieves the maximum value of 0.7±0.1 at $\Delta a/a=6.9\%$. However, when $\Delta a/a=9.7\%$, $S$ decreases to 0.6±0.1. The high value of $S$ obtained at low temperature of 573 K is attributed to an excessive number of misfit dislocations, resulting from the lattice mismatches between the Fe$_x$Pt$_{100-x}$ films and buffer layers, which are expected to promote the atomic diffusion. However, too large lattice mismatch leads to the decrease of $S$ for the Fe$_{52}$Pt$_{48}$ films[37].

![Fig.12 Degree of chemical ordering of Fe$_{38}$Pt$_{62}$ and Fe$_{52}$Pt$_{48}$ films expitaxially grown on buffer layers at 573 K as function of lattice mismatch between films and buffer layer, $\Delta a/a$, realized by changing materials of buffer layer[37]](image)

4.1.6 High pressure

Fig.13 shows the effect of annealing temperature on $S$ of the powders and bulk samples fabricated by warm compressing the initial powders under high pressure[38]. It is obvious that for the compacted samples, $S$ is of the order of zero before annealing temperature reaches 573 K, while it jumps to 0.9 at 673 K, indicating that $A1 \rightarrow L1_0$ transformation is almost accomplished. However, for the powder samples, 773 K is required to fulfill the ordering transformation. The high pressure is apparently responsible for the 100 K shift of the ordering temperature by compressing the fcc structure in the direction of $c$ axis to expedite the formation of the tetragonal one, $L1_0$.

4.1.7 External magnetic field

It is shown in Fig.14(a) that under the magnetic field of $1.03 \times 10^5$ A/m, the (001) and (110) superlattice peaks corresponding to the $L1_0$ phase for the FePt films, deposited on the Co$_{52}$Ag$_{48}$ underlayer, could be observed after annealing at 673 K; while without the magnetic field those typical peaks could no longer be observed until increasing the annealing temperature up to 773 K, indicating that the external magnetic field can induce the ordering of FePt films at reduced temperature.

![Fig.14 XRD patterns of FePt films grown on Co$_{52}$Ag$_{48}$ underlayer annealed at 673 K for 30 min with (a) and without (b) external magnetic field[27]](image)

4.1.8 Sputtering gas

It is exhibited in Fig.15 that the intensity ratio $I_{001}/I_{111}$ is utilized to describe the ordering, because the (001) peak corresponds to $L1_0$ phase, and (111) peak corresponds to $A1$ phase. $I_{001}/I_{111}$ for Kr-sputtering gas
develops from lower temperature of 523 K, while it develops from 573 K for Ne and 623 K for Ar. Meanwhile, $I_{001}/I_{111}$ at 673 K for Kr is 4.5, also distinctly higher than 0.55 and 0.75 for Ne and Ar, respectively. The reduced ordering temperature in the FePt films by the use of Kr as sputtering gas is interpreted to result from the fact that the Kr atoms possess the highest average kinetic energy, since the atomic mass of Kr is twice or four times heavier than that of Ar or Ne, respectively. Thus, the Kr sputtering gas can promote the ordering transformation[39].

With discussion above, we can conclude that the decrease of the ordering temperature is finally originated from the promotion of the atomic diffusion no matter what technical methods are employed.

4.2 Mechanism of magnetization and reversal magnetization of FePt based nano-composite films or particles

4.2.1 Effects of microstructure on coercivity of films or particles

Fig.16 exhibits the TEM images of the perfectly aligned (001) FePt films, epitaxially grown on the MgO (001) single crystal substrate at high temperature of $\geq 973$ K which is employed to ensure the high ordering of the films, for different nominal thickness, $t_N$(40–41).

It is observed in Fig.16 that interconnected network structures with voids in the shape of strip are observed for $t_N=60$ nm (Fig.16(a)). The voids expand with $t_N$ decreasing, and eventually some isolated particles emerge when $t_N=50$ nm (Fig.16(b)). It is notable that a remarkable change of the film takes place in the morphology from continuous network (Fig.16(b)) for $t_N=50$ nm to maze-like isolated particle for $t_N=45$ nm.
(Fig.16(c)). The size distribution of the particles is bimodal for the 20 nm-thick film (Fig.16(d)), consisting of larger particles with a typical size $d_2$ $>$ 200 nm and smaller ones in the size range from 10 to 50 nm. From experimental observation, the particle shapes are flat ellipsoids. On this situation, $a_h < a_e = a_s$, where $a_h$, $a_e$, $a_s$ are the half lengths of long axial, short axial and height of the ellipsoid, respectively. The size of the ellipsoidal particles $d_e$ is equal to $2a_h$. And in Fig.16(e) and Fig.16(f), the size of particles decreases from 20 $–$ 30 nm for $t_N=5$ nm, to 10 $–$ 20 nm for $t_N=3$ nm, then decreases to $<5$ nm when $t_N=1$ nm (Fig.16(g)). The whole size distribution is summarized in Table 1[23, 40–43].

### Table 1 Size distribution of particles in FePt films with different thickness [23, 40–43]

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<tr>
<th>$t_N$/nm</th>
<th>40</th>
<th>30</th>
<th>25</th>
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<tbody>
<tr>
<td>$d_e$/nm</td>
<td>$&gt;400$</td>
<td>10–50</td>
<td>10–20</td>
<td>$&gt;200$</td>
<td>50</td>
<td>20–30</td>
<td>10–20</td>
<td>$&lt;5$</td>
<td></td>
</tr>
</tbody>
</table>

It is displayed in Fig.17[40] that the initial magnetization curves, described to help analyzing the characteristic of the magnetization process, are of characteristic of domain wall displacement for the films with thickness $\geqslant 20$ nm since the magnetization easily progresses at low magnetic field (Curves (a), (b), (c) and (d)), indicating that those films consist of multi-domain continuous network or isolated maze-like particles, at least, isolated particles in bimodal distribution (Fig.16(d)). When $t_N$ decreases from 18 to 15 nm (from Curve (e) to (f)), the minor part of the magnetization rotation at large magnetic field corresponding to the single domain particle is observed. With $t_N$ further decreasing to 12 and 10 nm (Curves (g) and (h)), the fraction of the magnetization under large magnetic field increases, which is ascribed to the increased volume of single domain particles. Finally, the initial magnetization curves are of characteristic of magnetization rotation for $t_N < 8$ nm (Curves (i), (j) and (k)), indicating that these films are merely composed of single domain particles.

It is shown in Fig.18[40, 42] that $H_c$ increases appreciably with $t_N$ decreasing and holds a very low value when 50 nm $\leq t_N \leq 100$ nm. However, $H_c$ of 159.2 kA/m for $t_N=50$ nm jumps to 2 228.8 kA/m for $t_N=45$ nm, measured under a magnetic field of 5.5 T, with the morphology changing from continuous maze to particulate. A quite large $H_c$ value of about 3 422.8 kA/m for $t_N=40$ nm, measured under a magnetic field of 14 T, is achieved in spite of the fact that the domain wall displacement is dominant in magnetization process of the film. Measured under 14 T, $H_c$ increases with the decrease of $t_N$ when 5 nm $\leq t_N < 40$ nm; $H_c$ reaches the maximum value of 5 572 kA/m for $t_N=5$ nm; then begins to decrease when $t_N < 5$ nm; but still keeps a large value of 5 094.4 kA/m for $t_N=3$ nm; at last, it sharply decreases to 1 432.8 kA/m for the film with $t_N=1$ nm and $d_e < 5$ nm.

Since the $L1_0$ ordered FePt phase possesses large uniaxial $K_u$ in theory, it could be expected a huge $H_c$ value of 9 552 kA/m deduced from the coherent rotation model[41] ($2K_u/M_s \approx 9$ 552 kA/m) for ideally single domain $L1_0$ FePt particles. In practice, huge coercivity as high as 8 358 kA/m has been achieved at 4.5 K for the film of $t_N=5$ nm. This value is very approaching to the expected one.

#### 4.2.2 Domain wall pinning sites

It is remarkable that $H_c$ suddenly increases from 159.2 kA/m for $t_N=50$ nm to 2 288.8 kA/m for $t_N=45$ nm (Fig.18), with the morphology changing from continuous.
to particulate, resulting in the magnetic domain walls movement changing from full displacement to partial domain wall pinning due to the existence of void boundaries as domain wall pinning sites. However, no sharp increase is observed for the films deposited on SiO$_2$ substrate, although their morphology changes are the same as those of the films deposited on MgO (001) single crystal. Considering that the continuous polycrystalline films deposited on SiO$_2$ polycrystalline substrate have real grain boundaries, a high $H_c$ of $>1$ 034.8 kA/m is obtained in those kinds of continuous films due to strong domain wall pinning by the grain boundaries as well as by the twins within the grains, both of which are absent in the single crystalline films deposited on MgO (001) single crystal substrate[42]. KÜNDIG et al[44] have shown that a large $H_c$ of about 1 592 kA/m at the expense of magnetization can be obtained in melt-spun ribbons of a ternary Fe$_{35}$Pt$_{35}$P$_{30}$ eutectic composition by grain refinement as well as by introducing PtP$_2$ non-magnetic particles as possible pinning sites.

### 4.2.3 Critical size for single domain particle

The domain structure of L1$_0$ FePt particles changes from multi-domain to single domain when $d_s$ is below the critical size $D_{c1}$, approximately estimated[23] by using the following equation:

$$D_{c1} = \frac{24A_{m}K_{u}}{N_\perp M_s^2}$$

(3)

where $A_m=AS_{qs}/a_s$ is an intrinsic constant, $S_{qs}$ is the spin quantum number, and $N_\perp$ is the demagnetization factor along the height of the ellipsoid, given by[23]

$$N_\perp = 4\pi \left(1 - \frac{k}{(1-k^2)^{3/2}}\sin^{-1}\sqrt{1-k^2} + \frac{k^2}{1-k^2}\right)$$

(4)

$k=a_s/a_t$, $0<k<1$. By assuming $2a_s=t_N$, $2a_t=2d_s$, $k$, $N_\perp$, and $D_{c1}$ are estimated and calculated. $k$, $N_\perp$ and $D_{c1}$ are $0$–$0.2$, $7.81$–$12.56$, and $155$–$200$ nm, respectively.

From the magnetic force microscopy (MFM) observation, it has been revealed that in the case of $t_N=20$ nm, the calculated $D_{c1}$ of 155 nm is in good agreement with the experimental result of 180 nm. The slight difference is thought to result from the fact that the actual height of most particles is larger than the nominal thickness $t_N$[23].

The film in thickness of 40 nm is composed of particles of $d_s>400$ nm with multidomain structure. When $t_N$ decreases to 20 nm, it is comprised of two kinds of particles, the larger one about 200 nm with multidomain structure, the smaller one with single domain structure. The volume fraction of single domain particles increases with $t_N$ decreasing, leading to the increase of $H_c$ at first when $t_N \leq 40$ nm. This is mainly attributed to the change of the reversal magnetization process from magnetic wall displacement to wall pinning. With the decrease of the film thickness and the size of the particles, most of single domain particles appear in the film, accompanying with reversal magnetization model changing from wall pinning to magnetization incoherent rotation.

### 4.2.4 Critical size for reversal magnetization mode

In Fig.19, the FePt nano-particles in a carbon matrix are fabricated by depositing both FePt layers in thickness of 0.3–1 nm and carbon layers in thickness of 0.3–4.0 nm alternatively, followed by annealing at 873–1 073 K which leads to the formation of the nano-particles in carbon matrix. While the pure Fe$_{35}$Pt$_{35}$ nano-particles are synthesized by chemical method, considering the thermal...
Variation of coercivity with grain size (a) and degree of chemical ordering as function of inverse grain size (b) for FePt nano-particles in carbon matrix as function of C content in films[46−47].

CHRISTODOULIDES et al[45] have reported that the coercivity as a function of spherical particle size,
\[ d \]
follows the following relation:
\[ H_c = D_{c3} (\frac{d}{d_{c3}})^{3/2} \]
where \( D_{c3} \) is the superparamagnetic size and \( H_a (=2K_u/M_s) \) is the anisotropic field.

The \( H_c \) value of \(<796\) kA/m is relatively low (seen in Fig.19) even in the size of 10–20 nm. We believe that this is mainly attributed to the lower \( K_u \) value associated with incomplete atomic ordering since \( K_u \) is originated from the long-range ordering of alternatively stacked Fe and Pt layers along the c-axis. It is eventually affected by reduction in thickness along the c-axis to several atomic layers.

\( H_c \) decreases distinctly with decreasing particle size at \( d<8 \) nm, then vanishes at \( d=2 \) nm (Fig.19(a)). This is partly attributed to the increasing impact of the size effect, and the effect of chemical ordering on \( H_c \) is also supposed to be taken into account. TAKAHASHI et al [48] have found that there is a size dependence in the ordering process of FePt nano-particles. Although continuous FePt thin film is \( L1_0 \) perfectly ordered at 773 K, FePt nano-particles with size smaller than 5 nm cannot be ordered even annealing at 873 K for 1 h. Fig.19(b) shows that \( S \) decreases with reducing particle size. The results of both RONG et al[49] and CHRISTODOULIDES et al[45] in Fig.19(b) reveal a linear relation between \( S \) and \( 1/d \).

In theory, the minimal stable particle size, estimated from Ref.[43], is as small as 2.8 nm for \( L1_0 \) FePt at room temperature:
\[ D_{c3}=(60k_BT/K_u)^{1/3} \]

A detailed study on the ultrasmall nano-particles reveals that \( S \) sharply drops when \( d \) is below 3 nm[44] and FePt particles are not ordered to the \( L1_0 \) structure when \( d<2 \) nm, namely so called superparamagnetic limit[44, 46−47]. This agrees well with the experimental results. However, it is not clear about why the linear relation takes place. Further work is required to fully understand the relation among size effect, chemical ordering and the real effect of particle size on \( H_c \).

It is demonstrated that the coercivity is strongly associated with the characteristic size and the morphology of the assemblies of ferromagnetic particles. It is an effective way to induce a high \( H_c \) by controlling the size of the particles and by introducing domain wall pinning sites, i.e., nonmagnetic grain boundary and defects. The related studies on the relation of microstructures and coercivity are still going on.

4.2.6 Size effect on \( T_C \)

Fig.20 shows the dependence of \( T_C \) on the \( L1_0 \) FePt particle size. \( T_C \) decreases with particle size declining, remarkably when \( d<6 \) nm. For nano-particles smaller than 2 nm, \( T_C \) for the \( L1_0 \) phase cannot be generated since the phase transition cannot be realized. The shift of \( T_C \) from the value for bulk materials is expressed by[50]
\[ \frac{T_C(\infty)-T_C(d)}{T_C(\infty)} = \left(\frac{d}{d_0}\right)^{-1/\nu} \]
where \( T_c(d) \) is the Curie temperature as a function of \( d \), \( T_c(\infty) \) is the bulk Curie temperature, the value of \( d_0 \) is comparable to the lattice constant, and \( \nu \) is the critical exponent of the correlation length.

However, the effect of chemical ordering on the \( T_C \) should be considered in addition to the effect of particle size. This shows that the dependence of \( T_C \) and \( S \) on fixed particle size can be described in the region of \( 0.6 \leq S \leq 1 \) by a linear expression[47]:
\[ T_C = T_{c0} + a_0 S \]
where \( T_{c0} \) and \( a_0 \) are fitting parameters. It is notable that...
$T_c$ value in the Eq.(9) corresponds to the solid square in Fig.19(b), which is close to the real size effect on $T_c$ of fully-ordered $L1_0$ Fe$_5$Pt$_{48}$ nano-particles in Fig.20.

### References

7. GUO Peng-ju, LIU Xin-cai, PAN Jing, LI Yong, CUI Ping. Coercivity mechanism models and influential factors of Nd-Fe-B nano-composite magnets [J]. Rare Earth, 2009, to be submitted.

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