Hysteresis Loops, Critical Fields and Energy Products for Exchange-spring Hard/soft/hard Trilayers

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Macrosopic hysteresis loops and microscopic magnetic moment distributions have been determined by a three-dimensional (3D) model for exchange-coupled Sm-Co/α-Fe/Sm-Co trilayers with in-plane collinear easy axes. These results are carefully compared with the popular one-dimensional (1D) micromagnetic models and recent experimental data. It is found that the results obtained from the two methods match very well, especially for the remanence and coercivity, justifying the calculations. Both nucleation and coercive fields decrease monotonically as the soft layer thickness $L_s$ increases while the largest maximum energy product (roughly 50 MGOe) occurs when the thicknesses of hard and soft layers are 5 nm and 15 nm, respectively. Moreover, the calculated angular distributions in the thickness direction for the magnetic moments are similar. Nevertheless, the calculated nucleation and pinning fields as well as the energy products by 3D OOMMF are systematically smaller than those given by the 1D model, due mainly to the stray fields at the corners of the films. These demagnetization fields help the magnetic moments at the corners to deviate from the previous saturation state and facilitate the nucleation. Such an effect enhances as $L_s$ increases. When the thicknesses of hard and soft layers are 10 nm and 20 nm, respectively, the pinning field difference is as large as 30%, while the nucleation fields have opposite signs.

Keywords: hard/soft trilayers, hysteresis loops, energy products, micromagnetic calculation

1. Introduction

The concept of exchange-springs, i.e., exchange-coupled hard/soft composite nanomagnets, was proposed by Kneller et al. in 1991 [1]. In a typical composite magnet, the magnetizations of the soft phase respond to the applied field reversibly and therefore these magnets are called exchange-springs [1-3]. Two years later, Skomski and Coey predicted that the energy product of exchange-springs can be as large as 120 MGOe [4], which doubles that of the sintered Nd$_2$Fe$_{14}$B, the best performed permanent magnets up to now. Many efforts have been devoted to realize such a giant energy product in the past two decades [2-9]. Nevertheless, the experimental energy products are much smaller, even smaller than the sintered Nd$_2$Fe$_{14}$B magnets. Such a huge difference is called energy products in some literatures [10, 11]. Therefore, a reexamination of the present theories regarding to the hysteresis loops and energy products in the exchange spring materials is necessary.

Skomski and Coey made their prediction based on a one dimensional (1D) micromagnetic model [4], which was first utilized by Goto et al. in 1960s [12]. Similar model has been widely used in the past two decades, by Leineweber and Kronmüller et al. from Germany [13], by Fullerton and Jiang et al. from USA [2, 14], by Asti and Pellicelli et al. from Italy [15-17] and by Zhao et al. from China [10, 11, 18]. Such a simple model can reveal some important underlying physics with analytical formulas derived. However, it can not take account of the magnetic distributions in the film plane as well as the sophisticated microstructures existed in nature.

Experimentally, a major progress has been made in the last year [9], where an energy product of 60 MGOe has been achieved in NdFeB/FeCo multilayers by Cui et al. [9]. Such a high energy product surpasses that of the sintered Nd$_2$Fe$_{14}$B already. In the same time, the energy...
products in SmCo-Fe system have aroused intense interests and made progress. SmCo is the hardest magnetic material in the world which can supply the large coercivity whilst the soft Fe can provide high remanence. As early as 1995, SmCo/Fe/SmCo bilayers and multilayers have been fabricated by Al-Omari et al. [19], where the room-temperature energy product is only 6 MGOe. In 2002, Cheng et al. [20] fabricated the SmFeGaC/Fe permanent magnet, with the energy product increased to 8.2 MGOe. In 2005, Neu et al. [21] realized an energy product of 28.13 MG Oe (224 kJ/m$^3$) by improving the exchange coupling between SmCo and Fe layers. The Fe/Co mixture behavior enhances the interface exchange-coupling and improves the remanence and coercivity of the composite magnet [22]. Similar work has been done by Choi et al. [23], where the largest energy product increases to 32 MG Oe.

In this paper, the demagnetization process and hysteresis loops of SmCo/Fe/SmCo trilayer have been investigated using the 3D software (OOMMF) [25] and carefully compared with the results based on 1D analytic calculation.

### 2. Micromagnetic Model

The trilayer system studied in this paper is shown in Fig. 1, where a soft layer is sandwiched between two hard layers with collinear in-plane easy axes for both phases. A large magnetic field is applied along the easy axis direction and then gradually demagnetized to investigate the hysteresis loop related behaviors.

The three-dimensional (3D) calculation of the micromagnetic software package is based on the Landau-Lifshitz-Gilbert dynamic equation:

$$\frac{d\mathbf{M}}{dt} = - \gamma \mathbf{M} \times \mathbf{H}_{\text{eff}} - \frac{\gamma}{M_s} \mathbf{M} \times (\mathbf{M} \times \mathbf{H}_{\text{eff}}),$$

(1)

where $\mathbf{M}$ is the magnetization (A·m$^{-1}$), $\mathbf{H}_{\text{eff}}$ is the effective field (A·m$^{-1}$), $\gamma$ is the Landau-Lifshitz gyromagnetic ratio (m·A$^{-1}$·s$^{-1}$), and $\alpha$ is the dimensionless damping constant. The effective field is defined as follows:

$$\mathbf{H}_{\text{eff}} = - \mu_0 \frac{\partial E}{\partial \mathbf{M}},$$

(2)

The average energy density $E$ is a function of $\mathbf{M}$ specified by Brown’s equations [26, 27]:

$$E = A(r) \left[ \nabla M_z^2 \right] - K(r) \left( \frac{\mathbf{M} \cdot \mathbf{n}}{M_s} \right)^2 - \mu_0 \mathbf{M} \cdot \mathbf{H} - \frac{1}{2} \mu_0 \mathbf{H}_d(r) \cdot \mathbf{M},$$

(3)

where $A$ and $K$ are the exchange and anisotropy energy constants, respectively. $\mathbf{H}$ and $\mathbf{H}_d(r)$ are the applied and magnetostatic self-interaction fields while $M_s = |\mathbf{M}(r)|$ is the spontaneous magnetization. These equations hold for both the hard and soft phases. The four terms on the right side of Eq. (3) correspond to the exchange energy, the anisotropy energy, the applied field (Zeeman) energy, and the magnetostatic (demagnetization) energy, respectively.

The above 3D energy could be simplified to a one dimensional (1D) expression if we ignore the magnetostatic interactions, which are small for an infinitely large thin film with in-plane easy axis. In one dimension, the energy density per area in the film plane is [10, 28]:

$$v = \int_0^s \left[ A^1 \left( \frac{d\theta}{dz} \right)^2 + K^1 \sin^2 \theta - \mu_0 H_M \cos \theta \right] dz + \int_0^s \left[ A^2 \left( \frac{d\theta}{dz} \right)^2 + K^2 \sin^2 \theta - \mu_0 H_M \cos \theta \right] dz - \frac{2}{a} \frac{d^2}{dz^2} \left( \frac{m^2 - 1}{m^2} \right)$$

(4)

where $a$ is the distance between the adjacent atomic planes near the interface. $\theta$ is the angle between the magnetization and the applied field and $\mathbf{m}$ is the magnetization unit vector at the interface. For symmetry, only half of the trilayers has been included in the calculation. The three terms inside the bracket of the above formula are the exchange energy, the anisotropy energy and the Zeeman energy respectively, while the last term is the interface exchange coupling energy. A variational method [10, 28] is used to minimize the energy expressed in Eq. (4), which yields the equations for the angular distribution ($\theta$ as a function of $z$) as follows:

For $0 < z < L^h$:
where \( h = H/H_k \) represents the reduced applied field with \( H_k = 2K/(\mu_0 M_s) \) as the anisotropy field. \( \Delta = \pi A/\kappa \) is the Bloch wall width. \( \theta^h \) and \( \theta^s \) are the directions of the magnetization at the outer surfaces of the hard layer and the middle of the soft layer, respectively. From these equations the hysteresis loops and energy products can be calculated. The 1D calculation is basically an analytical one.

In this paper, Sm-Co is chosen as the hard layer while \( \alpha \)-Fe is the soft layer. The material parameters for the simulation are: \([2, 29]\) \( M'_h = 1.7 \times 10^5 \text{emu/cc}, M'_s = 0.55 \times 10^5 \text{emu/cc}, K' = 1000 \text{erg/cc}, K'' = 5 \times 10^3 \text{erg/cc}, A' = 2.8 \times 10^{-6} \text{erg/cm}, A'' = 1.2 \times 10^{-8} \text{erg/cm}, \) and \( A''_h = 1.8 \times 10^{-6} \text{erg/cm}. \) \( A, K, \) and \( M_s \) denote the exchange energy constant, the anisotropy constant, and the spontaneous magnetization, while the superscripts \( h \) and \( s \) refer to the hard and soft layers, respectively.

In the 3D simulation performed by OOMMF, the trilayer system is set as a tetragonal box, 300 nm \( \times \) 300 nm \( \times \) \( L_s \), where \( L_s \) is the thickness of the trilayer, which equals \( 2L'_h + L'_s \) according to Fig. 1. The thickness of the hard layer \( L'_h \) varies between 5 nm and 20 nm, while that of the soft layer \( L'_s \) varies between 2 nm and 20 nm. The system has been discretized into tetragonal cells with various sizes to find an optimum cell size where a good balance between the calculation precision and computational speed is obtained. The optimum cell size turns out to be 6 nm \( \times \) 6 nm \( \times \) 1 nm, \( i.e. \), with both length and width set as 6 nm while keeping a fixed thickness of 1 nm. The following calculations are based on such an optimum cell size unless otherwise specified. The applied field and the initial magnetization are along the x axis. Only the exchange interaction between the neighboring region pair is taken into account and the free boundary conditions are chosen.

### 3. Macroscopic Hysteresis Loops

Fig. 2 shows the macroscopic hysteresis loops of SmCo/Fe/SmCo trilayers with various soft layer thickness and a fixed hard layer thickness of 10 nm. The bottom and top panel correspond to the loops based on 3D and 1D methods respectively. The two methods give similar hysteresis loops and very close coercive fields, justifying the calculations. When \( L'_s = 2 \text{ nm} \), both models yield rectangular hysteresis loops with roughly the same nucleation and pinning fields. As \( L'_s \) increases, the squareness of the hysteresis loops becomes worse, accompanied by the gradual decrease of the nucleation and coercive fields. Here, the nucleation field is defined as the opposite of the applied field at which the magnetization has some obvious deviations from the previous saturation state. In the meantime, the difference between the two calculated loops based on the two models enlarges. One can see that when \( L'_s = 2 \text{ nm} \), the calculated difference for the pinning field is within 6%, which, however, increases to 30% as \( L'_s \) increases to 20 nm. The pinning field is defined as the opposite of the applied field at which the magnetizations in the hard layer reverse, which is displayed in the hysteresis loops as a great drop of the magnetization. Further, the nucleation process calculated by the 3D simulation is smoother than the one based on the 1D method. In the 3D simulation, the stray fields at the corner of the film help the nucleation process and thus the magnetization reversal. These stray fields, neglected in the 1D model, smooth the nucleation process and decrease the nucleation fields. This phenomena can be seen more

\[
\frac{L^h/2 + z}{\Delta^h/\pi} = \int_{\theta}^0 \frac{d \theta}{\sqrt{(\sin^2 \theta - \sin^2 \theta^h)^2 - 2h^2(\cos \theta - \cos \theta^h)}}.
\]  
\[
\text{For } -L^s/2 < z < 0:
\]

\[
\frac{L^h/2 + z}{\Delta^s/\pi} = \int_{\theta}^0 \frac{d \theta}{\sqrt{(\sin^2 \theta - \sin^2 \theta^s)^2 - 2h^2(\cos \theta - \cos \theta^s)}}.
\]

clearly from the in-plane magnetization distribution displayed in the next section.

Hysteresis loops have been calculated for SmCo \((L^h)\)/Fe\((L^s)\)/SmCo \((L^h)\) trilayers based on OOMMF for various \(L^h\), where \(L^h = 5\) nm (top), 15 nm (middle) and 20 nm (bottom) respectively. Similar to the case with \(L^h = 10\) nm. The squareness of the hysteresis loops is deteriorated as \(L^s\) increases, accompanied by the gradual increase of the remanence and the decrease of both nucleation and coercive fields.

4. Microscopic Magnetic Moment Distribution in the Film Plane

Fig. 4 shows the 2D evolution of the magnetizations with the applied field for a SmCo(10 nm)/Fe(20 nm)/SmCo(10 nm) trilayer. These in-plane magnetic distributions calculated by the 3D OOMMF, are located at the middle of the soft layer \((z = 0)\), which response to the applied field fast. At \(H = 14\) kOe, the magnetizations at the corners of the film deviates slightly from the applied field direction as shown in Fig. 4(a). However, most magnetizations still orient roughly at the applied field direction, demonstrating a state before the nucleation. The
The corresponding average magnetization of the system is 113 emu/cc, which is roughly the saturation magnetization of the system, as shown in Fig. 2. When the field decreases to 1 kOe (Fig. 4(b)), the magnetizations at the corners deviate more than 30º away from the previous saturation direction, whereas those at the center still orient at the saturation direction. This field corresponds to the nucleation field of the system. Such deviations will enhance and spread nearby through the exchange interaction upon the further decrease of the applied field. As the field decreases to ~3 kOe, obvious deviation can be observed at the top and bottom edges as shown in Fig. 4(c). In particular, the magnetizations at the corners deviate roughly 50º away from the the previous saturation direction already, whereas those at the center still orient at the previous saturation direction. As a result, the corresponding macroscopic magnetization is 107 emu/cc. Further decrease of the applied field will lead to the rotation of the magnetizations at the center. At $H = -6$ kOe (Fig. 4(d)), almost all magnetizations within this film plane are perpendicular to the previous saturation direction. Note that at this point, the response of the magnetizations at the edges with the applied field lag behind those at the centers, which have a deviation of roughly 60º only. This state is still some distance away from the coercivity of the system, as the magnetizations in the hard layer are still at the previous saturation state. The corresponding average magnetization is 512 emu/cc, as displayed in Fig. 2. The system will reach the coercive point at $H = -8.6$ kOe (Fig. 4(e)), where all the magnetizations in the middle plane of the soft layer orient at 120º away from the previous saturation direction. Further decrease of the applied field will lead to the saturation of the system. At $H = -13$ kOe, the trilayer saturates.

5. Angular Distribution at the Direction Perpendicular to the Film Plane and the Reversal Processes of Magnetization

The above magnetic moment distribution at the film plane can display nucleation modes very well, which starts at the surface of the soft layer. However, the important magnetic reversal processes through domain wall motion is not revealed. To do this, the angular distribution at the $z$ direction should be discussed.

Fig. 5 shows the angular distributions of the magnetization at the thickness direction for the SmCo(10 nm)/Fe(5 nm)/SmCo(10 nm) trilayers based on 1D and 3D models. For the 1D calculation, the angular distribution at the $z$ direction could be obtained by minimizing the energy density expressed in Eq. (4), which is coherent at x-y plane. For the 3D calculation, the angular distribution is obtained by averaging the component of all the magnetic moments in the applied field direction with a certain value of $z$. Due to the symmetry, only magnetization distributions in the half of the trilayers have been drawn and shown.

According to the 3D calculation, a small deviation of the magnetization from the previous saturation state occurs at $H = -24$ kOe, which is called nucleation according to Brown. At this field, a 9.2º infant domain wall has been formed with $\theta^h = 0^\circ$, $\theta^s = 7.8^\circ$ and $\theta^0 = 9.2^\circ$. Only about 15% of the domain wall is within the soft phase due to the very thin soft layer. The angular increase of the magnetization can be observed as the applied field decreases. Exception is observed at the surface of hard phase, where $\theta^0$ keeps a constant of 0º from nucleation to pinning. Due to the large thickness of the hard phase, the angle at the surface of the hard phase $\theta^0$ obeys the Stoner-Wohlfarth (SW) model [30], where $\theta^0 = 0^\circ$ before the magnetic reversal and $\theta^0 = 180^\circ$ after the reversal. In the meantime, the domain wall is pushed toward the hard phase gradually. When $H = -28$ kOe, $\theta^0$ and $\theta^h$ increase to $33.7^\circ$ and $39.2^\circ$ respectively, where 86% of the domain wall locates at the hard phase. When the applied field decreases to the coercivity ($H = -30.51$ kOe), $\theta^0$ and $\theta^h$ increase to $60.9^\circ$ and $69.5^\circ$ respectively.

The 1D calculation gives similar angular distributions, where the nucleation occurs at a more negative applied field ($H = -27.5$ kOe). At the nucleation point, a 43.5º domain has been formed in the trilayers, with 21%
located at the soft phase. The corresponding $\theta^0$ and $\theta'$ are $34.4^\circ$ and $43.5^\circ$ respectively. A slight further decrease of the field will lead to a large increase of both $\theta^0$ and $\theta'$. In the meantime, the domain wall moves toward the hard phase gradually. At $H = -28.2$ kOe, $\theta^0$ and $\theta'$ increase to $80.8^\circ$ and $95.7^\circ$ respectively, where roughly 85% of the domain wall is at the hard phase. At the coercivity (-29.8 kOe), $\theta^0$ and $\theta'$ increase to $121.7^\circ$ and $107.9^\circ$ respectively, where only 11% of the domain wall is in the soft phase. Further decrease of the applied field leads to the irreversible reversal of the magnetizations of the whole system.

Although the results of the 1D and 3D are similar, there are some differences between them. The angles according to the 3D calculations are systematically smaller than those based on the 1D calculations. In addition, the motion of the domain walls from the soft to the hard phase is more evident based on the 1D calculation, where the percentage of the domain wall within the hard phase increases from 79% at the nucleation to 89% at the coercivity. More importantly, the nucleation process based on the 3D calculations is much slower, leaving a large gap between nucleation and coercivity.

The angular distributions for other soft layer thicknesses have also been calculated and compared. Fig. 6 shows the calculated angular distributions for SmCo (10 nm)/Fe (10 nm)/SmCo (10 nm) trilayers based on the two methods, which are similar to those for $L' = 5$ nm. Noticeably, the angles according to the 3D calculations are systematically smaller than those based on the 1D calculations. Moreover, the motion of the domain walls from the soft to the hard phase is evidenced in the 1D results, where the percentage of the domain wall within the hard phase rises from 60% at the nucleation to 76% at the coercivity.

Compared with the case with $L' = 5$ nm, the percentage of the domain wall within the soft phase is larger, accompanied by the drop of both nucleation and coercive fields. According to the 3D simulation, a domain wall of $7.4^\circ$ forms in the trilayers at the nucleation point with 68% in the hard phase. The angle of the magnetization increases with the decreasing applied field. When $H$ decreases to $-13$ kOe, the domain wall develops to $45.8^\circ$ and increase to $76.0^\circ$ at the coercive point, with 69% and 71% in the hard phase respectively.

In general, the 3D calculation yields better results than those given by the 1D calculation, as can be seen from Figs. 2 and 3 that the calculated coercivity and energy products based on the 3D calculation is generally smaller than that from the 1D calculation. For example, the calculated coercivity decreases from 15.3 kOe to 8.6 kOe as $L'$ increase from 10 nm to 20 nm for SmCo(10 nm)/Fe/SmCo(10 nm) trilayers according to the 3D calculation (see Table 1). On the other hand, the corresponding coercivity reduces from 18.2 kOe to 8.6 kOe according to the 3D calculation (see Table 1). Similar coercivity can be obtained for other $L'$ as can be seen from Table 2 that the coercivity is not sensitive to the hard layer thickness. Whilst experimental coercivity drops from roughly 10 kOe to 7.5 kOe achieved by Sawatzki et al. for SmCo(5 nm)/Fe(25 nm)/SmCo(25 nm) trilayers. Therefore, the

![Fig. 6. (Color online) Calculated spatial distribution of magnetic orientation $q$ in the thickness direction at various applied fields for SmCo(10 nm)/Fe(10 nm)/SmCo(10 nm) trilayers. (a) Calculated by the 3D simulation and (b) calculated by the 1D model. Due to the symmetry, only magnetization distributions in the half of the trilayers has been drawn and shown.](image)

**Table 1.** Calculated magnetic properties for SmCo(10 nm)/Fe($L'$)/SmCo(10 nm) trilayers based on OOMMF (3D) and the simplified model (1D), where $H_N$, $H_C$, and $H_P$ denote the nucleation, coercive and pinning fields respectively, while $(BH)_{max}$ and $M_r$ stand for the maximum energy product and the remanence.

<table>
<thead>
<tr>
<th>$L'$ (nm)</th>
<th>Method</th>
<th>$H_N$ (kOe)</th>
<th>$H_C$ (kOe)</th>
<th>$H_P$ (kOe)</th>
<th>$(BH)_{max}$ (MG Oe)</th>
<th>$M_r$ ($\times 10^3$ emu/cc)</th>
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<td>2</td>
<td>1D</td>
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<td>67.3</td>
<td>67.3</td>
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<tr>
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<td>63.5</td>
<td>16.9</td>
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<td>15.3</td>
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coercivity gap between the experiment and the theory is shortened according to the 3D calculation, which can include the local stray fields. Similar situation holds for the energy product gap. However, the 3D calculation requires much more computational power than the 1D calculation so that the mesh size in the 3D calculation is larger than that for the 1D calculation. Therefore it is hard to say which model is better. We use both methods in this paper and compare the results to justify our methods.

### Table 2. Calculated magnetic properties for SmCo($L^h$)/Fe($L^s$)/SmCo($L^h$) trilayers based on OOMMF, where $H_N$, $H_C$, and $H_P$ denote the nucleation, coercive and pinning fields respectively, while $(BH)_{\text{max}}$ and $M_r$ stand for the maximum energy product and the remanence.

<table>
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<th>$L^s$ (nm)</th>
<th>$L^h$ (nm)</th>
<th>$H_N$ (kOe)</th>
<th>$H_C$ (kOe)</th>
<th>$H_P$ (kOe)</th>
<th>$(BH)_{\text{max}}$ (MG Oe)</th>
<th>$M_r$ ($\times 10^3$ emu/cc)</th>
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</table>

### 6. Critical Fields and Energy Products

From above discussions, it becomes clear that the most important magnetic properties for hard/soft multilayers are critical fields, *i.e.*, nucleation and coercive fields. Fig. 7 highlights the calculated critical fields based on OOMMF and the 1D model for SmCo(10 nm)/Fe($L^s$)/SmCo(10 nm) trilayers, as functions of the soft layer thickness. It can be seen that the two models yield similar critical fields. For small $L^s$, the coercivity equals the nucleation field, whereas $H_N$ is larger than $H_C$ for large $L^s$. As the soft layer thickness increases, both nucleation and coercive fields decrease. In the meantime, the gap between the nucleation and coercivity rises. Similar trend has been obtained for other hard/soft bilayers [10, 15, 18]. Experimentally, Sawatzki et al. [24] found that the coercivity is equal to the nucleation field in SmCo$_5$(25 nm)/Fe/SmCo$_5$(25 nm) trilayers for small $L^s$, whereas $H_C$ becomes to deviate from the nucleation field when the soft layer thickness is above a critical value of 8.8 nm. The gap between $H_N$ and $H_C$ also increases with $L^s$. However, the experimental critical thickness is much larger than those shown in Fig. 8.

![Fig. 7.](image_url) Calculated nucleation and coercive fields as functions of the soft layer thickness based on OOMMF (dotted lines) and 1D analytical model (Solid lines).

![Fig. 8.](image_url) Calculated second-quadrant energy product $(BH)_{\text{max}}$ for SmCo(10 nm)/Fe($L^s$)/SmCo(10 nm) trilayers. (a) Energy products based on OOMMF; (b) energy products based on the 1D calculation.
zation fields, taken into account by the OOMMF, facilitate the nucleation and lead to considerably lower nucleation fields. The stray fields will also have some influence on the calculated coercivities and energy products, which, however, is less significant compared with the former. For example, when $L' = 20$ nm, the calculated nucleation fields have opposite signs, whereas the difference for the coercivities are within 1%. Therefore, the gap between the nucleation and the coercivity is larger according to the 3D calculation.

Fig. 8 shows the Calculated energy products for SmCo($10$ nm)/Fe($L'$)/SmCo($10$ nm) trilayers based on 1D and 3D methods. Both models give similar energy products. In particular, the calculated maximum energy products are very close, justifying our calculation. For $L' \leq 10$ nm, both models give exactly the same $(BH)_{\text{max}}$. Here the $(BH)_{\text{max}}$ is achieved at the points before the nucleation takes place and thus not affected by the nucleation difference caused by the different models. For $L' \geq 15$ nm, the $(BH)_{\text{max}}$ based on the 3D calculation are smaller, due to the smaller nucleation fields obtained. However, the difference between the calculated $(BH)_{\text{max}}$ according to the two models are still within 10%.

Importantly, both models yield the largest $(BH)_{\text{max}}$ at $L' = 15$ nm, which is around 40 MGOe. These results agree well with available experimental data. Experimentally, Sawatzki et al. [24] obtained an energy product of 40 MGOe in SmCo/$s$/Fe/SmCo$_{5}$ trilayers at $L' = 12.6$ nm.

The calculated magnetic properties for SmCo($10$ nm)/Fe($L'$)/SmCo($10$ nm) are summarized in Table 1. One can see that both models yield similar results. In particular, the calculated remanence, coercivity and energy products based on the two methods are very close, where the difference is within 10%. As $L'$ increases, the critical fields go up monotonically whereas the remanence falls down. Therefore, the energy products have a peak at $L' = 15$ nm.

On the other hand, the nucleation and pinning fields based on OOMMF calculations are systematically smaller than those based on the 1D model. The difference enlarges as $L'$ increases. When $L' = 2$ nm, both differences for the nucleation and pinning fields are within 10%. When $L' = 20$ nm, however, the pinning field difference is as large as 30%, while the nucleation fields have opposite signs. It should be addressed that although the nucleation field difference is significant at large $L'$, it does not affect much on the energy products or the hysteresis loops. As has been mentioned above, the nucleation based on the 3D is a slow and smooth process. The magnetizations do not change much after the nucleation. The small nucleation fields compensated by the slow process from the nucleation to pinning, leading to roughly the same coercivity as that based on the 1D model. The coercivity mechanism is pure pinning for small $L'$ according to the OOMMF calculation. It changes to a hybrid one at large $L'$, where the coercivity is in between the nucleation and pinning fields. The coercivity mechanism based on the 1D model is more complicated. The coercivity equals both nucleation and pinning fields at small $L'$. Such a mechanism changes to a pure pinning one at intermediate $L'$ and finally to a hybrid one at large $L'$.

In the particular case for $L' = 15$ nm (see Table 1), in both models the $H_N$ are quite different. However, the coercivities is very similar as the coercivity is not determined by the nucleation field in this system. Rather, it is completely determined by the pinning field according to the 3D calculations. In the 1D calculation, the situation is slightly different; $H_c$ is in between the nucleation and the pinning fields but is more close to the pinning field. Further, the two models suggest similar $M_s$ and $H_c$, which are important reference properties for the largest energy product. Nevertheless, the $(BH)_{\text{max}}$ values are more different as $(BH)_{\text{max}}$ is not determined by these two values completely. $(BH)_{\text{max}}$ is also closely related to the shape of the hysteresis loop in the second quadrant. In particular, the nucleation field based on the 3D model is much smaller than that according to the 1D model, leading to a smaller $(BH)_{\text{max}}$.

The calculated magnetic properties corresponding to Fig. 3 for SmCo($L^h$)/Fe($L'$)/SmCo($L^h$) with other three values of $L^h$ are summarized in Table 2. One can see that the largest energy products also occur at $L' = 15$ nm for all values of $L^h$. The highest $(BH)_{\text{max}}$ rises from 27.5 MGOe to 52.8 MGOe as $L^h$ decreases from 20 nm to 5 nm. Experimentaly, energy products of 40 MGOe has been achieved by Sawatzki et al. [25] for SmCo$_{5}$(25 nm)~/Fe(12.6 nm)/SmCo$_{5}$(25 nm) trilayers. Both the value of $(BH)_{\text{max}}$ and the soft layer thickness at which the highest energy products occur are in good agreement with the present model. On the other hand, the hard layer thickness at which the largest $(BH)_{\text{max}}$ occurs is about 5 nm, which is much smaller than the experimental one. Similar results have been obtained by other groups [4, 15-17]. For example, Skomski and Coey [4] recommend that the hard layer thickness to be less than 3 nm for the giant energy product to be realized. These deviations come from the fact that the effective anisotropy of the hard phase in the experiment might be less than the theoretical one [10]. Further, some possible distributions of easy axis orientations (although very small in the experiments) might affect the energy product and shift the optimum hard layer thickness to a higher value [11]. The formation of the Fe-Co [23,
7. Conclusions

In this study, the properties of Sm-Co/α-Fe/Sm-Co tri-layers have been studied using a 3D micromagnetic software, OOMMF and compared carefully with those given by the 1D analytical model. It is found that the calculated hysteresis loops and angular distributions are consistent with each other, ensuring the reliability of our calculations. In particular, both nucleation and coercive fields decrease as the soft layer thickness increases. The difference for the remanence, coercivity and maximum energy products based on different models are less than 10%. The calculated largest $(BH)_{\text{max}}$ is around 50 MGOe, achieved at $L_s = 15$ nm ($L_h = 5$ nm), which is consistent with available experimental data.

The calculated nucleation and pinning fields based on OOMMF are systematically smaller than those according to the 1D model, where the difference increases as $L_s$ increases. These differences are due to the large stray fields acting on the magnetic moments at the corners of the film planes, which is ignored in the 1D calculation.

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