Coercivity of a percolative magnetic system

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A model that describes the behavior of the coercivity $H_c$, of coalescing Ni films, as a percolation phenomenon is presented. We show that there is a nonmonotonic dependence of the coercive field as a function of the occupation probability $p$. This behavior is a function of cluster size and topology, and is independent of the growth direction of the thin film. Based on these ideas we predict theoretically, and confirm experimentally, the dependence of $H_c$ as a function of external magnetic field orientation. For a [111] oriented fcc film, if $p$ is less than the percolation threshold $p_{cr}$, a sixfold symmetry is observed. However, if $p = p_{cr}$, the percolation is isotropic.

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The concept of percolation was introduced in the realm of science by Broadbent and Hammersley1 in 1957. Since then, it has been applied to many problems in completely different fields, ranging from natural sciences to sociological phenomena.2 In physics, percolation has been useful in the study of microscopic3 as well as macroscopic phenomena.4,5

In particular, granular systems inherently prone to percolation, have experimentally shown magnetic anomalies when the grains are magnetic. Panina et al.6 found that the effective magnetic permeability of composite materials containing small Fe particles (of $1-2 \ \mu \text{m size}$) tends to zero near the percolation threshold. Xiao and Chien7 observed that across the whole volume fraction range, in granular Fe-(SiO$_2$) solids, magnetic coercivity experiences dramatic variations due to the change of grain size and percolation effects. In thin films the relation between magnetism and percolation also has been investigated: for example, the transition from superparamagnetism to ferromagnetism,8 while recently Gor'kov and Krein9 studied the transition from paramagnetic to conducting ferromagnetic phases of manganites by percolation theory. Moreover, Choi et al.10 observed that the coercivity $H_c$ of Ni films, whose magnetic and structural properties are found in the literature,11 exhibits a nonmonotonic behavior, related to topological morphology changes, which coincides with the onset of percolation. However, no theoretical ideas have been advanced which directly connect coercivity and percolation.

In order to study changes in $H_c$ as a function of morphology in Ni, Choi et al.10 performed a well controlled experiment evaporating Ni films, by molecular beam epitaxy (MBE), on top of various thickness $t_{Cu}$ copper buffer layers. The morphology of the Cu buffer layer changes with $t_{Cu}$. First isolated copper clusters form, but as $t_{Cu}$ increases pairs of clusters coalesce. As more Cu is deposited a critical thickness is reached and the Ni covered Cu clusters percolate. In order to assure that all changes in the coercivity of Ni are due only to changes in morphology, all other structural parameters of the epitaxial Ni were kept constant. These parameters include: the thickness (100 Å), the level of contamination, the growth parameters, the capping layer width (150 Å of Al), the (111) crystalline orientation perpendicular to the substrate, and the mosaic spread. All these structural and chemical properties were checked using in situ Auger electron spectroscopy (AES), reflection high energy electron diffraction (RHEED), low energy electron diffraction (LEED) and ex situ x-ray diffraction (XRD), scanning electron microscopy (SEM), and atomic force microscopy (AFM).

Two major variations of the coercivity $H_c$ become apparent as the morphology reaches the percolation threshold; the magnitude of $H_c$ changes in a nonmonotonic fashion and the in-plane angular anisotropy changes as well. Both of them display a distinct feature, which coincides with the crossing of the percolation threshold. Our theoretical model explains these two distinct features, that is: (i) the magnitude of $H_c$ as a function of the copper substrate thickness; and (ii) the dependence of $H_c$ on the direction of the applied field $H$. The model shows that $H_{cr}$, as a function of site occupation probability $p$, has a nonmonotonic dependence due to the fractional variation of the cluster sizes and their topology, as they form during the film deposition process, and it is also shows that this behavior is independent of the growth direction of the magnetic film. Moreover, also the symmetry of $H_{cr}$, when $p = p_{cr}$, is in quantitative agreement with experiment.

Our model focuses upon the morphological changes that take place during the growth of a thin film, which in its early stage is characterized by a homogeneous distribution of the small islands. When additional material is deposited these islands grow in size, due to the large mobility of the small droplets, until coalescence sufficient for percolation is achieved. Based on this the coercivity $H_{cr}(p)$, as a function of occupation probability $p$, can be understood assuming that the magnetization reversal mechanism depends on cluster size and topology: (i) if the particles are smaller than a certain critical size $z_{cr}$, of the order of a domain wall, $H_c$ is large, due to the fact that for such particles wall nucleation is energetically unfavorable. Consequently, the magnetization...
reversal proceeds, for instance, through coherent rotation of the spins, \(^{12}\) (ii) for simply connected particles, whose sizes are greater than \(z_{cr}\), the coercive field \(H_c\) is smaller than the former since now a domain wall \(^{11}\) can nucleate. Thus, the magnetization reversal is achieved by processes that require less energy, predominantly by domain wall nucleation and displacement; and (iii) larger clusters may be multiply connected and the domain walls might get pinned by the voids, especially if these are of considerable size. Thus, when the fraction of voids is large enough the inversion mechanism depends both on cluster size and topology, since domain wall formation is likely, but their propagation is hindered by pinning to the voids that are present.

As usual \(H_c\) is defined as the reversal field required to achieve zero magnetization. To determine the coercivity we introduce the reduced magnetization \(M_{\mu}(H)\), which corresponds to the quotient between the actual and the saturation magnetization of a single cluster. The indices \(\mu=s, b, v\) label clusters of the type (i), (ii), and (iii), as described above. Thus, the analytic expression for the film magnetization \(M(H)\) is

\[
M(H) = \rho_s M_s(H) + \rho_v M_v(H) + (1-\rho_s-\rho_v) M_b(H),
\]

where \(\rho_{\mu}\) is the fractional area occupied by each type of clusters. Consequently we obtain the coercivity \(H_c\) solving the equation \(M(H_c)=0\).

The fractional areas \(\rho_s\) and \(\rho_v\), of the different type clusters, are obtained by using the Monte Carlo technique. \(^{14}\) In this algorithm each of the \(L\) sites of a lattice is occupied by a particle with probability \(p\). This produces a configuration of \(Lp\) particles. A cluster is defined as follows: two occupied sites belong to same cluster if they are nearest neighbors. The calculation is iterated keeping all the parameters (system size and occupation probability) fixed and changing the configuration the particles adopt. The final values of \(\rho_s\) and \(\rho_v\) are obtained averaging over a number of random configurations. Convergence is obtained after \(10^3\) Monte Carlo steps and averaging over ten random configuration.

We now formulate a criterion to classify a cluster as belonging to type (iii) as defined above. Designating as \(f\) the ratio between the number of empty sites inside a cluster \(\eta_v\) and the number of occupied sites in the same cluster \(\eta_o\), that is \(f=\eta_v/\eta_o\), we classify the cluster as belonging to class (iii) if \(f>f_{cr}\).

Moreover we make the following simple assumption: that only two types of islands are important, as far as magnetic properties are concerned. Clusters of type (i) and (ii), with broader loops, switch by magnetization rotation, while the larger type (ii) ones, with narrower loops, only switch after they nucleate a domain wall. To obtain \(H_c\) as a function of \(p\) we need to know the form of \(M_s(H)=M_s(H)\) and \(M_b(H)\) in the region \(H=0\) to \(H_b\), and we denote as \(H_b\) and \(H_c\) the broad and narrow loop values, respectively. The approximations we use to fit the experimental relations between magnetization and applied field \(M_{\mu}(H)\) are

\[
M_{\mu}(H) = M_r \tanh\left(\frac{H+H_{\mu}}{2q_{\mu}}\right),
\]

where \(M_r\) is the reduced remanent magnetization, \(\mu=s\) and \(b\), and \(q_{\mu}^{-1}\) is proportional to the slope of the corresponding hysteresis loop at \(M_{\mu}=0\).

The results of implementing the model outlined above are given in Fig. 1. As expected, the fractional area of small clusters \(\rho_s\approx1\) for \(p<<1\), and decreases to \(\rho_s\approx0\) as \(p\) grows. On the other hand, the fractional area \(\rho_v\approx0\) for \(p<<1\) and grows very fast when the percolation threshold is approached. Finally, \(\rho_v\) undergoes an abrupt drop when the dominant contribution is due to the formation of large clusters without voids. The kinks of \(\rho_v\) arise from finite-size effects. Replacing the numerical data obtained for \(\rho_s\) and \(\rho_v\), and combining Eq. (2) with \(M(H_c)=0\), we obtain the coercivity as a function of \(p\) as displayed in the insert of Fig. 1(a).

In order to make further progress an assumption has to be made on the unknown relation between the experimentally measured \(^{10}\) copper layer thickness \(t_{Cu}\) and \(p\). We choose the simplest alternative: a linear relation. The two values required to determine this fit are chosen as follows: (i) the minimum of \(H_c\); and (ii) the value for which \(H_c\) versus \(p\), in the insert of Fig. 1, ceases to be flat as a function of \(t_{Cu}\). This yields

\[
t_{Cu}[^{\AA}] = 780p + 650.
\]

The above fit is certainly not unique, but depends on the value we adopt for \(z_{cr}\). Consequently, this fit is valid only for \(650< t_{Cu}[^{\AA}] \leq 1430\). However, the results for \(H_c\) we obtain for \(650< t_{Cu}[^{\AA}] \) are insensitive to this choice and
thus we refrain from postulating a more sophisticated fit, which would imply additional parameters.

On the basis of the experiments we adopt $M_r=0.69$ for both switching mechanisms, since it was observed that $M_r$ does not depend on $\theta_C$. For the slopes we adopt $q_s=q_p=50$ (Oe), and stress the fact that the main features of $H_c=H_c(p)$ do not depend on the detailed form of $M_s(H)$ and $M_b(H)$. In fact, we found no significant changes in the behavior of $H_c$ for two extreme cases: (a) a rectangular hysteresis loop, and (b) assuming linear fits of $M_s(H)$ [$M_b(H)$] which go through $H_c$, $H_s$ and $M_r$. Always (i) if $p < p_c$, then $H_c$ versus $p$ is flat, since a single switching mechanism dominates and thus $H_c=H_s$; (ii) independent of the form of $M_s(H)$ and $M_b(H)$ a minimum is obtained around the percolation threshold $p_c$; and, (iii) a maximum of $H_c$ appears for $p > p_c$, which decays to the minimum value of $H_c$ as $p$ grows. This behavior is insensitive to the form of $M_s(H)$ and $M_b(H)$. We employed the value $f_c=0.5$ in the insert Fig. 1(a); changing the latter only narrows or enhances the width of the $p > p_c$ maximum. The fact that the $H_c$ versus $\theta_C$, displayed in the insert, is similar both for triangular and square lattices is a strong indication that this behavior is universal, i.e., does not depend on the growth direction of the thin film.

On the other hand, the angular dependence of the coercivity is also modified by percolation. For the small clusters we consider a single domain thin film, with its surface oriented in the [111] direction. In spherical coordinates the direction of the magnetization $\vec{M}$ is defined by the angles $\theta$ and $\phi$ that $\vec{M}$ makes with the Cartesian axes ($z$ axis is oriented along the [111] direction, while the $x$ axis is parallel to the [$\bar{1}10$] direction). The film parameters are defined as follows: $K$ is an effective anisotropy constant and $M_0$ is the saturation magnetization. The external field $H$ is confined to the $x,y$ plane, and $\phi_H$ is the angle between $H$ and $x$.

Three terms contribute to the total energy (per unit volume) $E_T$: the magnetocrystalline anisotropy $E_K$, the Zeeman term $E_Z$ and the magnetostatic energy $E_M$. $E_K$ determines the easy magnetization directions, which lie along the diagonals of the cubic cell (in the [111], [111], [111], and [111] directions). In this coordinate system

$$E_K = -\frac{\sqrt{2}}{3} K \sin^3 \theta \cos \theta \sin (3 \phi) + \frac{K}{12} \cos^2 \theta \left(6 - 7 \cos^2 \theta\right),$$

$$E_Z = -H M_0 \sin \theta \cos (\phi - \theta_H),$$

$$E_M = -2 \pi N M_0^2 \sin^2 \theta,$$

where $N$ is the demagnetizing factor. Defining the dimensionless energy $\varepsilon = 3E_T/K$, we can write

$$\varepsilon = \epsilon_1 - h \sin \theta \cos (\phi - \theta_H),$$

where $\epsilon_1 = 3(E_k + E_M)/K$ and $h = 3M_0 H/K$. For equilibrium it is required that $\partial \varepsilon / \partial \theta = 0$ and $\partial \varepsilon / \partial \phi = 0$. Stable (unstable) equilibrium is attained if the determinant of the associated Hessian matrix is positive (negative). As the (negative) field intensity $H$ is increased, $M$ rotates first gradually and then abruptly towards the unstable field direction. Analytically this is fulfilled when one of the following conditions is satisfied: $\partial^2 \varepsilon / \partial \theta^2 = 0$ or $\partial^2 \varepsilon / \partial \phi^2 = 0$.

The dimensionless field $h$ can thus be evaluated. However, only the lowest negative eigenvalue obtained for $h$ is physically meaningful; we make sure that this corresponds to a crossover from a minimum to a maximum by checking the sign of $\partial^2 \varepsilon / \partial \phi^2$. Figure 2 compares the results of these calculations with experiment; the error bars of the latter are smaller than the size of the dots in the figure. The parameters used in the computations are: $M_b=500$ G, $K = 2022$ erg/cm$^3$, and the demagnetization factor $N = 0.05$. Assuming a Gaussian distribution of the anisotropy axes of magnetic clusters around the mean orientation, it is found analytically that $K$ is reduced in good agreement with the fits obtained from Fig. 2.

$H_c$ has a sixfold symmetry, related to the fact that we are dealing with the (111) face. However, an additional condition is also required: that the spins adopt a small out of the (111) plane component, since otherwise the sixfold symmetry is quenched. Thus, as the system is rotated around the (111) axis [perpendicular to the (111) plane] the spins are barely drawn towards the easy [111] axis, all in satisfactory agreement with experiment.

On the other hand, when the percolation threshold $p_c$ is reached, the system behavior is dominated by the large clusters. In that case, the dipolar interaction becomes larger and has the effect of aligning in plane the total magnetization of the cluster. Thus the anisotropy energy $E_k(\theta = \pi/2) = 0$ is independent of $\phi$, which was verified experimentally as shown in Fig. 2. A very small orientation dependence of the

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**FIG. 2.** Coercivity $H_c$ as a function of the magnetic field direction $\phi_H$. The continuous line is the theoretical sixfold fit of the experimental data depicted, for $p < p_c$, by the full dots. The full squares illustrate the isotropic dependence of $H_c$ versus $\phi_H$ for $p = p_c$, which corresponds to $\theta_C = 100$ nm, measured at room temperature. The experimental error is smaller than the size of the full dots and squares.
The decrease of $H_c$ are small, and thus the magnetization, by nucleation and motion of domain walls. This forms. We have also shown that, when a fcc film is grown in continuous film value, since as $p$ grows a single large cluster forms. We have also shown that, when a fcc film is grown in the [111] orientation, a sixfold symmetry of $H_c$ for $p < p_{cr}$ exists, which crosses over to an isotropic behavior for $p = p_{cr}$.

In conclusion, we have developed a model which displays coercivity and provides an understanding for the relation between co-

Below the percolation threshold $p_{cr}$ most of the clusters are small, and thus the $H_c$ versus $t_{c,r}$ relation in Fig. 1 is flat. The decrease of $H_c$ is related to the increase in the number of large clusters, until they exceed the number of small ones, allowing a crossover from magnetization rotation to domain-wall nucleation and motion, to be the dominant magnetization switching mechanism. However, there is an increase of $H_c$, in the vicinity of $p_{cr}$, since inside the clusters the presence of internal nearest neighbors empty sites pin the domain-wall displacement. Finally, $H_c$ decreases up to its continuous film value, since as $p$ grows a single large cluster forms. We have also shown that, when a fcc film is grown in the [111] orientation, a sixfold symmetry of $H_c$ for $p < p_{cr}$ exists, which crosses over to an isotropic behavior for $p = p_{cr}$.

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16 $E_x = \frac{1}{2} \kappa \cos(6(\theta - \theta_0))$, where $\kappa$ is the bulk anisotropy constant. If the axis direction $\theta_0$ is assumed to have a Gaussian distribution of width $\Delta$, then $\langle E_x \rangle = \frac{1}{2} \kappa \exp(-9\Delta^2)\cos(6\theta)$. For $\Delta \approx 25^\circ$ this yields $K \approx \kappa/10$.
17 Experimentally, the $\theta_0$ dependence of the coercive field $H_c$ was obtained at room temperature on a vibrating sample magnetometer. The values were obtained from full $M-H$ loops that were taken at $2^\circ$ intervals, for in-plane angles from zero through $180^\circ$ or $360^\circ$, depending upon the sample. All samples exhibited sixfold symmetry below $p_{cr}$, similar to that displayed in Fig. 2.