Magnetic patterning: local manipulation of the intergranular exchange coupling via grain boundary engineering

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Magnetic patterning, with designed spatial profile of the desired magnetic properties, has been a rising challenge for developing magnetic devices at nanoscale. Most existing methods rely on locally modifying magnetic anisotropy energy or saturation magnetization, and thus post stringent constraints on the adaptability in diverse applications. We propose an alternative route for magnetic patterning: by manipulating the local intergranular exchange coupling to tune lateral magnetic properties. As demonstration, the grain boundary structure of Co/Pt multilayers is engineered by thermal treatment, where the stress state of the multilayers and thus the intergranular exchange coupling can be modified. With Ag passivation layers on top of the Co/Pt multilayers, we can hinder the stress relaxation and grain boundary modification. Combining the pre-patterned Ag passivation layer with thermal treatment, we can design spatial variations of the magnetic properties by tuning the intergranular exchange coupling, which diversifies the magnetic patterning process and extends its feasibility for varieties of new devices.

Information technology makes great use of various magnetic devices including magnetic patterned media1, magnetic random access memory2, magnetic domain wall devices3,4, and spin-wave devices5. A common approach to fabricate these nanostructure devices is the conventional lithography technique6,7 – excess regions are physically removed to define the desired spatial patterns with uniform magnetic properties. On the other hand, if the magnetic properties can be spatially controlled without physical etching, it can open up a new route for device designs with novel functionality. The so-called magnetic patterning is proposed to create the spatial distribution of magnetic properties by rearranging interface1, inducing order-disorder phase transformation8–10, doping additional elements11, or generating chemical reactions12,13. The magnetic patterning is now mainly realized by ion implantation14, and several unique devices have been demonstrated, such as the one-way domain wall motion shift registers15 or magnonic crystals16. Other magnetic patterning methods with different working mechanisms, including nano-indentation17, local thermal treatment18, or local anisotropy modification by adjacent layers19,20, have also been proposed. Most of the methods share the similar strategy by altering the spatial distribution of the effective magnetic anisotropy (K_{eff}), sometimes accompanying the change of saturation magnetization (M_s) as in ion implantation. To escape from the constraints imposed by changing magnetic

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anisotropy and/or saturation magnetization, is it possible to find another tuning knob to realize magnetic patterning for device designs?

In addition to modifying $K_{\text{eff}}$ and $M_s$, controlling microstructure is another way to modify magnetic properties, especially with regard to grain boundaries which has been widely investigated in recording media$^{21}$ and permanent magnets$^{22}$. By adding intergranular additives, one can engineer grain boundaries in recording media and permanent magnets for magnetic hardening, which may accompany modified intergranular exchange coupling. However, these approaches usually alter grain boundary structure uniformly in the entire sample and no spatial manipulation can be achieved.

In this work, we demonstrate a different approach for magnetic patterning by locally engineering the grain boundaries with altered intergranular exchange coupling. With pre-patterned passivation layers and thermal treatments, we demonstrate the magnetic patterning on Co/Pt multilayers (MLs), which are used as the model system due to its easy fabrication and perpendicular anisotropy for potential applications$^{23}$. The magnetic analyses and microstructure investigations are carried out to reveal the working principles for this magnetic patterning method. With rapid thermal annealing (RTA) process, we found that stress relaxation in Co/Pt MLs brings about modification of grain boundary structures and the corresponding magnetic properties. Meanwhile, by adopting pre-patterned passivation layers, we can spatially control the degree of stress relaxation in the designated regions with lateral modulation of the nucleation field ($H_n$), the domain wall propagating field ($H_p$), and the domain structures in Co/Pt MLs. In addition, we find that the same magnetic patterning can be achieved by Joule heating. By applying electric current, the magnetic properties in the specific regions can be altered. Different from the reported magnetic patterning methods, our unique grain boundary engineering approach alters local magnetic properties leaving both $K_{\text{eff}}$ and $M_s$ intact. Our proposed approach opens up a new avenue for magnetic patterning and can be easily incorporated into the existing process for device fabrication, for example, the BEOL (back-end-of-line) process, how the magnetic devices are usually integrated with other components$^{24,25}$.

**Results and Discussions**

**Magnetic patterning processes.** The flowchart of the proposed method for magnetic patterning is shown in Fig. 1(a). First, 20 nm thick Ag lines with different line width are pre-patterned on Co/Pt MLs by e-beam lithography and lift-off. Before RTA, the whole sample shows uniform magnetic properties.
The sample is then annealed by using RTA in vacuum (5 × 10⁻⁵ Torr) with duration of 20 seconds. Regions with or without Ag capping show different responses to RTA, leading to heterogeneous distributions of the magnetic properties in different spatial regions. After RTA, the remaining Ag patterns can be selectively etched away by ammonia-based etching solution (NH₃OH·H₂O₂ = 1:1) without damages to the magnetic properties of Co/Pt MLs (see supplementary Figure 1), yielding a flat magnetic film with spatially manipulated magnetic properties.

The hysteresis loops of the magnetic patterned Co/Pt MLs after 350°C annealing is measured by focused magneto-optical Kerr effect (FMOKE) with a 2 μm probing spot in diameter, as shown in Fig. 1(b). Both loops probed in regions with or without the Ag capping layer, denoted as Ag-capped and uncapped respectively, show sharp magnetization transitions. However, the magnetic field (~2000 Oe) for magnetization switching in the uncapped regions is larger than that (~1500 Oe) in the Ag-capped regions.

The atomic force microscopy (AFM) and the magnetic force microscopy (MFM) images of the patterned Co/Pt MLs are taken before the removal of the Ag capping as shown in Fig. 1(c). In the left panel of Fig. 1(c), the AFM and MFM images respectively show the surface topography and magnetization distribution of Co/Pt MLs, after MLs are partially saturated, that is, an applied magnetic field with the magnitude between the switching field of uncapped and Ag-capped regions is applied on MLs before MFM images are taken. The AFM and MFM images clearly reveal that the regions with bright contrast in AFM images, corresponding to Ag-capped regions, possess reversed magnetization, showing bright contrast in MFM images; the regions with dark contrast correspond to the uncapped regions without magnetization reversal. On the other hand, the AFM and MFM images of the ac-demagnetized state are shown in the right two panels of Fig. 1(c). The regions with or without the Ag capping layer reveal quite different domain sizes: the Ag-capped regions show the larger domain sizes, but the uncapped regions show smaller ones. Our results clearly demonstrate that local manipulation of the magnetic properties and the ac-demagnetized domain structure in Co/Pt MLs can be realized by combining Ag pre-patterns and the RTA process.

**Working principles of the magnetic patterning: magnetic analyses.** Because the key ingredients for our proposed magnetic patterning are RTA and Ag capping layers, to understand the underlying mechanism, we first evaluate the modifications of magnetic properties on Co/Pt MLs with varied RTA temperatures and capping conditions. The representative out-of-plane hysteresis loops and initial curves of Co/Pt MLs with or without 20 nm Ag capping layer are shown in Fig. 2(a). All hysteresis loops of as-deposited and RTA samples reveal sharp magnetization transition, also observed in the magnetically patterned Co/Pt MLs shown in Fig. 1(b). It implies that the magnetization reversal for both sheet and patterned Co/Pt films is dominated by nucleation of reversed domains, followed by rapid domain wall expansion[26]. Based on the hysteresis loops and initial curves, we can extract the nucleation field (Hₙ, the field beyond which the first magnetization reversal occurs from the saturation state) and the domain wall propagating field (Hₚ, the field required for the onset of domain wall depinning). Both fields increase with elevating RTA temperature (T_ann) below 350°C, as plotted in Fig. 2(b) (see Supplementary Figure 2 for the details when T_ann is above 350°C). We also observe that Hₙ is always smaller than the corresponding Hₚ and the Hₚ/Hₙ ratios for all samples remain constant, as shown in Fig. 2(c). It is worth mentioning that the samples capped with Ag layers reveal smaller Hₙ and Hₚ than those without Ag under the same RTA condition, consistent with the case found in the corresponding regions in the Ag-patterned samples.

The magnetization reversal behavior of Co/Pt MLs, including Hₙ and Hₚ, is reported to be determined by the interfacial anisotropy (Kᵥ) originating from Co/Pt interfaces[27]. The effective magnetic anisotropy (K_eff) of Co/Pt MLs with perpendicular magnetic anisotropy can be expressed by

$$K_{eff} = 2K_S/t + K_V = M_SH_K/2$$

where t is the thickness of each Co layer, Kᵥ the bulk anisotropy, and H_K the anisotropy field. After annealing, K_eff of Co/Pt MLs has been reported either to be reduced due to intermixing with lower Kᵥ[27] or enhanced by interfacial alloying of ordered CoPt with higher Kᵥ[28]. Therefore, both Hₙ and Hₚ can change when K_eff is altered. However, as shown in Fig. 2(d), both the Hₙ and Mₛ are independent of T_ann here.

It suggests that the significant enhancement of Hₙ and Hₚ after RTA cannot be simply explained by the modification of K_eff – a different mechanism is in order to explain the observed variations of Hₙ and Hₚ.

In addition to the changes of Hₙ and Hₚ, we also observed that sizes of the ac-demagnetized domains (D_domain) are shrunk by elevating T_ann, as shown in Fig. 3(a). Furthermore, the Ag-capped samples show the less modification of D_domain compared to the uncapped ones. Based on the micromagnetic simulations reported by R. H. Victora et al[28], with K_eff and Mₛ almost constant, the ac-demagnetized domain size of Co/Pt MLs can be determined by intergranular exchange coupling, which can be evaluated by using the ΔM curves[29]. In Fig. 3(b), the upper and lower panels show the ΔM curves of the uncapped and the Ag-capped samples, respectively. All samples exhibit positive ΔM peaks, indicating an exchange interaction dominated magnetization reversal[30]. The higher positive maximum intensity of the ΔM curve (ΔM(H)_max) indicates the stronger intergranular exchange coupling. For all samples, the intergranular exchange coupling reduces with the elevating T_ann as the ΔM(H) max drops. The Ag-capped samples show a smaller drop of ΔM(H) max with T_ann than that of the uncapped counterparts, implying that the...
extra Ag capping layer hinders the reduction of intergranular exchange coupling during the RTA process. Furthermore, we also notice that the samples with smaller intergranular exchange coupling (lower ΔM(H)Max and smaller D\text{domain}) shows higher H\text{n}, indicating that the intergranular exchange coupling should be an important factor for the magnetization reversal of Co/Pt MLs here.

To quantitatively evaluate the influence of intergranular exchange coupling on magnetization reversal, we checked the effective activation volume (V\text{act}) of each sample and used it as an indicator for the strength of intergranular exchange coupling. Note that V\text{act} represents the size of an effective magnetic volume composed of physical grains coupled together through intergranular exchange coupling\textsuperscript{33,34}. The effective activation V\text{act} can be obtained by extracting the characteristic half time (t\text{1/2}) from magnetization relaxation curves of different applied magnetic fields (H\text{app}) and fitting curves into the equation shown below\textsuperscript{26},

$$\ln(t_{1/2}) = V_{\text{act}}M_S/k_B T (H_C - H_{\text{app}})$$  \hspace{1cm} (2)

where k\textsubscript{B} is the Boltzmann constant, T is measurement temperature (ambient temperature in this case), and H\textsubscript{C} is the coercivity, respectively. As shown in Fig. 3(c), both V\text{act} and D\text{domain} decrease with increasing T\text{ann}, indicating that the RTA process reduces intergranular exchange coupling. Besides, because the D\text{domain} and V\text{act} are both affected by intergranular exchange coupling\textsuperscript{33,34}, the linear dependence between V\text{act} and D\text{domain} revealed in Fig. 3(d) finds its natural explanation. Moreover, we found that the H\text{n} is proportional to the reciprocal of the activation volume (1/V\text{act}), as plotted in Fig. 3(e), showing the increase of H\text{n} accompanied by the reduction of V\text{act}. This relationship can be explained by the curling mode for nucleation\textsuperscript{35}. In the curling mode, nucleation occurs via forming a curled spin structure with a lower energy barrier compared to that of nucleation via the Stoner-Wohlfarth model with coherent rotation. The nucleation energy barrier of the curling mode increases with shrinking V\text{act} because it costs more
exchange energy to form the curled spin structure. Therefore, the smaller $V_{\text{act}}$ is, the higher $H_n$ becomes (see Supplementary Note 1 for details). Noticeably, although the changes of $1/V_{\text{act}}$ for the Ag-capped and the uncapped samples are different, indicated by the length of the blue and red dash lines in Fig. 3(e) respectively, all data points of these two types of samples follow the same linear dependence. It indicates that $H_n$ is directly determined by $V_{\text{act}}$ of the Co/Pt MLs. The role of the Ag capping layer is mainly to control the size of $V_{\text{act}}$, that is, the intergranular exchange coupling.

Now, we would like to discuss the effect of the intergranular exchange coupling on $H_p$. When intergranular exchange coupling is lowered at grain boundaries, grain boundaries behave effectively as domain wall pinning sites so domain wall pinning strength can be enhanced. Therefore, reduction of the intergranular exchange coupling leads to a smaller $V_{\text{act}}$ as well as a higher domain wall pinning site density ($n_{\text{pin}}$) simultaneously. The parameter $1/V_{\text{act}}$, a function of intergranular exchange coupling here, can also be used as an indicator for $n_{\text{pin}}$. As the model proposed by V. Zabolotskii et al., higher $n_{\text{pin}}$ leads to higher $H_p$. Hence, $H_p$ increases with increasing $1/V_{\text{act}}$ and decreasing intergranular exchange coupling, as $H_n$ does. Consequently, the direct dependence of $H_p$ and $H_n$ on the intergranular exchange coupling explains the correlation between $H_p$ and $H_n$ shown in Fig. 2(c).

**Working principles of the magnetic patterning: structural and microstructural analyses.** To reveal how intergranular exchange coupling changes with RTA processing and to identify what the role of Ag passivation is, we investigate the structural changes occurred during RTA by performing 0–20 X-ray
diffraction (XRD). Figure 4(a) shows the variations of the central \( (n=0) \) and satellite \( (n=-2, -1, 1) \) peaks of the uncapped (Ag-capped) samples with increasing \( T_{\text{ann}} \) are shown in the upper panel (lower panel). (b) Dependence between the reciprocal of activation volume \( (1/V_{\text{act}}) \) and the change of strain \( (-\Delta \varepsilon_z) \). (c) STEM-HAADF images and corresponding STEM-EELS mapping of Co L\(_{2,3}\) edge are shown for the as-deposited and 350 °C annealed uncapped samples. The red dashlines on HAADF images indicate the position of grain boundary; The yellow dash-lines on the Co mapping point out the offset of Co layers between adjacent grains. (d) Co line scans of the 350°C annealed sample extracted from the Co mapping (the same image shown in (c), but rotated by 90°) on different positions: the green line shows the Co distribution across the whole grain, and the red line is along the grain boundary (as indicated on the Co mapping image). The blue circles on the Co line scan along the grain boundary indicate the distribution of out-diffusion Co via grain boundary creep.

Figure 4. Structural and microstructural analyses of the Co/Pt MLs with different capping conditions (a) 0–2θ X-ray diffraction patterns of Co/Pt MLs with different \( T_{\text{ann}} \). The central \( (n=0) \) and satellite \( (n=-2, -1, 1) \) peaks of the uncapped (Ag-capped) samples with increasing \( T_{\text{ann}} \) are shown in the upper panel (lower panel). (b) Dependence between the reciprocal of activation volume \( (1/V_{\text{act}}) \) and the change of strain \( (-\Delta \varepsilon_z) \). (c) STEM-HAADF images and corresponding STEM-EELS mapping of Co L\(_{2,3}\) edge are shown for the as-deposited and 350 °C annealed uncapped samples. The red dashlines on HAADF images indicate the position of grain boundary; The yellow dash-lines on the Co mapping point out the offset of Co layers between adjacent grains. (d) Co line scans of the 350°C annealed sample extracted from the Co mapping (the same image shown in (c), but rotated by 90°) on different positions: the green line shows the Co distribution across the whole grain, and the red line is along the grain boundary (as indicated on the Co mapping image). The blue circles on the Co line scan along the grain boundary indicate the distribution of out-diffusion Co via grain boundary creep.

diffraction (XRD). Figure 4(a) shows the variations of the central \( (n=0) \) and the satellite \( (n=-2, -1, 1) \) XRD peaks of Co/Pt MLs (111) and (222) planes with \( T_{\text{ann}} \). The central peak position is determined by the average d-spacing of Co/Pt MLs, and the satellite peak intensity reflects interface sharpness between Co and Pt\(^{38}\). All samples show a central peak shift to a higher 2θ angle with elevating \( T_{\text{ann}} \) accompanying the conserved satellite peak intensity when \( T_{\text{ann}} \) is below 350°C. The conserved satellite peak intensities indicate a conserved layered structure; therefore, the observed central peak shift should not be related to the bulk diffusion between layers, for example, intermixing or alloying, which would destroy the layered structure. On the other hand, by wafer curvature measurements\(^{39}\), we confirm that stress relaxation occurred after RTA process. The as-deposited samples reveal the presence of the in-plane bi-axial compressive stress but the annealed samples show less compressive stress. (see Supplementary Table 1 for
on the Co L 2,3-edge EELS signal difference between the layers, we found that the Co layered structure was interrupted by additional Pt atoms, resulting in reduced intergranular exchange coupling 44. Broken, bringing about Co deficient grain boundaries, where the direct exchange coupling between Co atoms is reduced after annealing) versus the corresponding 1/V act, we observe a linear dependence between −Δε z and 1/V act as observed in the H n vs. 1/V act, implying the stress relaxation process may strongly correlate with the intergranular exchange coupling of Co/Pt MLs.

To clarify the correlation between stress relaxation and intergranular exchange coupling, we performed cross-sectional imaging with high-angle annular dark field (HAADF) as well as electron energy loss spectra (EELS) of Co L 2,3-edge mapping by using scanning transmission electron microscope (STEM). As shown in Fig. 4(c), the columnar grains and layered structure of Co/Pt MLs are clearly shown in the STEM-HAADF images for the as-deposited (left) and the 350 °C annealed samples (right). The different contrast of the adjacent grains originates from different crystal orientations of grains with respect to the electron beam direction and the brighter contrast inside each grain represents Pt layers with higher atomic number compared to the darker Co layers. Based on the contrast between grains and the lattice images, we can distinguish the grain boundaries, marked by red dash lines on the HAADF images. We subsequently performed STEM-EELS Co L 2,3-edge mapping on the same region, where the bright contrast represents the Co distribution. For the as-deposited sample, the Co layers are continuous between the adjacent grains, i.e. across the corresponding grain boundary. On the other hand, for the 350 °C annealed sample, although the Co layers are still distinguishable and continuous inside each grain, the Co layered structure is out-of-sync at grain boundaries, offsetting to each other across grain boundaries, as marked by the yellow dash-line in the 350 °C annealed Co mapping image. As shown in Fig. 4(d), we also extract the line scan data through the grain (shown by the green dash-line) and along the grain boundary (the red dash-line) from the Co L 2,3-edge mapping of 350 °C annealed sample. Based on the Co L 2,3-edge EELS signal difference between the layers, we found that the Co layered structure after 350 °C RTA keeps clear periodic distribution of chemical composition throughout the grain, but becomes blurred along the grain boundary. That is, the EELS signal maximums are lower and the distributions of Co along the film normal are wider at grain boundary (red) compared to the case through the grain (green). Moreover, based on the line scan along the grain boundary, we also observed that some Co diffused out of the layered structure along grain boundary, circled by blue in Fig. 4(d), which was not found in the as-deposited samples. (see Supplementary Figure 3 for more details)

To understand the modification of microstructure at grain boundaries and its corresponding changes of the magnetic properties, the stress relaxation process during RTA will be discussed first. Because of the deposition stress during sputtering40 (revealed in wafer curvature measurements of the as-deposited sample), and the thermal stress41 during RTA, an in-plane bi-axial compressive stress exists in Co/Pt MLs. When temperature exceeds a critical value, diffusion is activated to rearrange the atoms to release the accumulated stress, the so-called diffusion creep, which has been believed as one of the mechanisms governing the stress relaxation in metal films when temperature is raised42. Because the layered structure inside grains is conserved but becomes blurred for the chemical composition at grain boundary, we propose that the stress relaxation during RTA is mainly due to diffusion creep via grain boundary diffusion, i.e. the so-called Coble creep.43 (see Supplementary Figure 4 for the possibility of stress relaxation via grain growth) That is, an out-diffusion mass flow via grain boundaries occurs during RTA processing, including both Co and Pt atoms, which accompanies stress relaxation. Both the out-diffusion Co along grain boundaries and the chemical composition blurred grain boundaries as observed in the EELS Co mapping image are the evidences for the Coble creep mechanism for stress relaxation. Meanwhile, because the out diffusion mass flow along grain boundary includes both Co and Pt atoms, the composition of grain boundaries can be disturbed and the continuity of Co across grain boundaries can also be broken, bringing about Co deficient grain boundaries, where the direct exchange coupling between Co atoms was interrupted by additional Pt atoms, resulting in reduced intergranular exchange coupling 44. Besides, the out-of-sync grain boundaries with offset Co layers may also decrease the intergranular exchange coupling by reducing the direct Co-Co exchange coupling between the adjacent grains 44. Consequently, the larger −Δε z occurred in Co/Pt MLs after RTA process indicates that larger amount of mass flow has been driven to modify grain boundaries, leading to a lower intergranular exchange coupling strength and reduced V act in Co/Pt MLs.

In addition, it has been reported (mostly in BEOL process) that when the stress relaxation is dominated by diffusion creep, an additional passivation layer, possessing good adhesion and low interdiffusivity with the adjacent metal film, can significantly hinder the stress relaxation of metal films by changing its diffusion kinetics 41,43. That is, the free surface of the metal film with a lower energy barrier for diffusion is replaced by the interface between the passivation layer and the metal film with a higher energy barrier. The energy barrier for diffusion raises significantly and impedes the diffusion of atoms or vacancies, and therefore hinders the diffusion creep. Thus, a proper passivation layer suppresses the stress relaxation during annealing, for example the AlO x on Al 46 and the SiN x on Cu41. In our case, the Ag capping layer has been reported to offer a robust Ag/Pt interface with good adhesion and only monolayer interdiffusion 47 so the Ag capping layer is selected to suppress the stress relaxation. We confirm that no significant
compositional profile changes of Ag and Pt occur during RTA (see Supplementary Figure 5). We also exclude the possibility that Ag diffusion modifies magnetic properties (see Supplementary Note 3). As shown in Fig. 4(a), under the same $T_{\text{ann}}$, the $2\theta$ shift of the uncapped samples is always larger than those of the Ag-capped samples, implying that an extra Ag capping layer indeed suppresses the stress relaxation during RTA. With a Ag capping layer, the fast free surface diffusion is blocked by the robust Ag/Pt interface and the grain boundary diffusion is therefore retarded, leading to less stress relaxation and the less amount of modified grain boundaries. Furthermore, we found that the trend of $-\Delta \varepsilon_z$ vs. $1/V_{\text{act}}$ lies in the same line for both the uncapped and Ag-capped samples, shown in Fig. 4(b), indicating that the intergranular exchange coupling of Co/Pt MLs is strongly related to the degree of stress relaxation, which can be altered by either RTA temperature or passivation layer. This concept can be further verified by using different passivation layers. For example, Ta is another good passivation layer due to good adhesion between Ta and Pt, but SiO$_2$ is not (see Supplementary Figure 6 for details). Simply by selecting proper pre-patterned passivation layers, we can alter the diffusion kinetics and laterally modify the intergranular exchange coupling to achieve the proposed magnetic patterning.

**Magnetic patterning by local Joule heating.** As discussed in previous paragraphs, the magnetic properties can be locally manipulated by controlling stress relaxation during annealing process. However, the infrared lamp used in our RTA process, providing uniform heating area for the whole sample, should not be the only driving force for stress relaxation. Here we demonstrate how Joule heating generated by currents can replace the RTA process so that we can selectively anneal the designated magnetic wires. The Joule heating method provides another route to manipulate the spatial distribution of magnetic properties. As shown in Fig. 5(a), by applying current pulses (40 pulses with the current density of 45MA/cm$^2$ and the duration of 50 ms for each pulse) into the selected magnetic wires, only the wires with current are annealed. The same passivation effects as RTA are observed by using Joule heating annealing. As shown in Fig. 5(b), the hysteresis loops acquired by FMOKE reveal that the magnetization switching of the Ag-capped regions occurs at a lower field than that of the uncapped regions. Inset of Fig. 5(b) shows the partially reversed MFM images of the magnetic patterned magnetic wire at the partially saturated states. The Ag-capped region is indicated by blue dot-squares with different width of Ag pre-pattern (4 and 2 $\mu$m, respectively).

**Uniqueness of proposed magnetic patterning.** To diversify the magnetic patterning methods is of great importance for designing new devices and extending their functionality. The most unique feature for our magnetic patterning method is to adjust intergranular exchange coupling without significantly changing $K_{\text{eff}}$ and $M_s$ in all regions, which have not been achieved before and can provide a new degree of freedom to design magnetic devices. The modified intergranular exchange coupling has been shown to alter the $H_n$ and $H_p$ due to the modified $V_{\text{act}}$ and $n_{\text{pin}}$, respectively. In current-driven domain wall motion devices, the magnetic patterning has been used for creating the nucleation sites or pinning sites in magnetic wires to control the domain wall motion. Our magnetic patterning method, which alters the $H_n$ and $H_p$ in local regions, can be used for defining the storage bits in the magnetic wires for the so-called...
racion. Besides, we can generate different $H_n$ laterally in the magnetic wires. Therefore, the magnonic crystal can be potentially obtained by introducing various $H_n$ instead of $M_s$ in different area to achieve periodical magnetization states (partially switched state, like the cases in Fig. 1(c) and the insertion of 5(b)).

Compared to ion implantation or other magnetic patterning methods, our approach can achieve a nanoscale magnetic patterning as demonstrated previously and is possible for a large-scale patterning region (across the whole wafer) without special or expensive equipment. Besides, different from ion implantation that may damage the dielectric layers and the logic devices underneath within the VLSI (very-large-scale integration) framework, one of the advantages of our method is BEOL compatible (as discussed in Supplementary Figure 6). Furthermore, by Joule heating annealing, our magnetic patterning cannot only be performed on the selected magnetic wires, but can be potentially carried out after the whole IC packaging process, opening a new route for post-production programing of magnetic devices. Furthermore, because the passivation effect on stress relaxation is a common phenomenon observed in many other metal films, we believe our approach can be further extended to other multilayers structure, for instance Co/Pd$_{49}$, Co/Ni$_{50}$ multilayers, or even other kinds of perpendicular magnetic anisotropy materials. By properly choosing a passivation material, we can achieve magnetic patterning in other systems.

Regarding the resolution of our magnetic patterning, our proposed method is mainly confined by the roughness of the domain wall. The domain wall is defined by the edges of Ag wires because the regions with Ag capping show small $H_n$ and $H_p$. Since we used a lift-off process to define the Ag lines, we may introduce artificial defects at edges and thus induce the domain wall roughness. If we optimize the lift-off process or use the etching process to well define the edges of Ag wires, the resolution of our proposed method can be further improved.

Conclusions

We have demonstrated a new magnetic patterning method by adopting pre-patterned passivation layers, Ag, with thermal treatments, including RTA and Joule heating annealing. The working principle of our proposed magnetic patterning relies on the control of stress relaxation in Co/Pt MLs during thermal treatment. The modification of grain boundaries occurs during thermal treatments due to Coble creep, leading to the changes of intergranular exchange coupling. The pre-patterned Ag passivation layer is shown to control the degree of stress relaxation and thus the strength of intergranular exchange coupling. When the intergranular exchange coupling is altered, the $H_n$, $H_p$, and $D_{\text{domain}}$ are modified accordingly; therefore, the magnetic properties in the regions can be engineered depending on with Ag or without Ag capping, leading to lateral distribution of magnetic properties. Our proposed approach diversifies the magnetic patterning methods, that is, changing $H_n$ and $H_p$ without altering $M_s$ or $K_{\text{eff}}$ which may offer a new degree of freedom for designing novel magnetic devices.

Methods

We prepared the Co/Pt MLs by a ultra-high vacuum magnetron sputtering system with the base pressure about $4 \times 10^{-8}$ Torr. The multilayers structure we used in this work is substrate/Ta(3 nm)/Pt(7 nm)/[Co(0.4 nm)/Pt(2 nm)]$_8$, deposited on thermally oxidized Si (100) substrates. A 20 nm thick Ag passivation layer was then deposited on some of the multilayers. The working pressure of Ar was 7 mTorr for Pt, and 3 mTorr for the others. Rapid thermal annealing with infrared lamps was subsequently performed at temperatures from 200°C to 350°C with a duration of 20 seconds and a ramp rate of 25°C/sec under the vacuum of $5 \times 10^{-5}$ Torr.

To demonstrate the magnetic patterning, we first pre-patterned the Ag capping layer with different stripes width by e-beam lithography and lift-off process. After the Ag pattern was prepared, RTA was carried out. After RTA, the Ag stripes were selectively removed by NH$_3$OH(30wt%):H$_2$O$_2$(30wt%) etching solution, which effectively etches Ag away without reacting with the MLs. For the magnetic patterning on a single magnetic wire by Joule heating annealing, we use conventional photolithography and ICP-RIE (inductively coupled plasma reactive ion etching) for defining wires of Co/Pt MLs; the Ag pre-patterns are subsequently defined by photolithography and lift-off. By controlling the amplitude of current density, the numbers of pulses, and the duration for each pulse, we can modify magnetic properties. Here, we use 40 pulses with the current density of 45MA/cm$^2$ and the duration of 50 ms for each pulse. By in-situ monitoring the resistance change with external magnetic fields, that is, measuring magnon magnetoresistance$^{31}$, we can obtain the coercivity so that we can adjust the current pulses and durations accordingly to modify the magnetic properties.

A vibration sample magnetometer (VSM) was used for in-plane and out-of-plane hysteresis loops of the Co/Pt MLs. $H_n$, $M_s$, and $H_p$ were extracted from the out-of-plane hysteresis loops with initial curves, where $H_n$ is defined by the onset of magnetization drop from saturated state in the hysteresis loops, and $H_p$ is defined by the first slope change of initial curves; $H_K$ was estimated from the saturation field of in-plane loops. VSM was used to measure magnetization relaxation curves for calculating the effective activation volume and to acquire $\Delta M$ curves. The magnetic force microscopy (MFM) and corresponding atomic force microscopy (AFM) were carried out for estimating the ac-demagnetized domain size and revealing magnetic patterns in local regions. The ac-demagnetized domain size ($D_{\text{domain}}$) was quantitatively estimated by using two dimensional fast Fourier transformations (2D-FFT) technique for each
corresponding MFM image. The focused magneto-optical Kerr effect measurement with 2μm laser spot was performed for the hysteresis loops in local regions. The wafer curvature measurement was carried out for stress analyses in Co/Pt MLs. The 0–20 X-ray diffraction (XRD) was performed for measuring the change of strain of Co/Pt MLs before and after RTA process. The cross-sectional high angle annular dark field images and corresponding element mapping were acquired by Cs-corrected scanning transmission electron microscope (STEM) and electron energy loss spectra (EELS), respectively, with the electron microscope operating at 200kV.

References
1. Chappert, C. et al. Planar patterned magnetic media obtained by ion irradiation. Science 280, 1919–1922 (1998).
2. Ikeda, S. et al. A perpendicular-aniostropy CoFeB-MgO magnetic tunnel junction. Nature materials 9, 721–724 (2010).
3. Allwood, D. A. et al. Magnetic domain-wall logic. Science 309, 1688–1692 (2005).
4. Parkin, S. S., Hayashi, M. & Thomas, L. Magnetic domain-wall racetrack memory. Science 320, 190–194 (2008).
5. Neusser, S. & Grundler, D. Magnonics: Spin Waves on the Nanoscale. Adv Mater 21, 2927–2932 (2009).
6. Geissler, M & Xia, Y. N. Patterning: Principles and some new developments. Adv Mater 16, 1249–1269 (2004).
7. Terris, B. D. & Thomson, T. Nanofabricated and self-assembled magnetic structures as data storage media. J Phys D-Appl Phys 38, R199–R222 (2005).
8. Lai, C. H., Yang, C. H. & Chiang, C. C. Ion-irradiation-induced direct ordering of L1(0) FePt phase. Appl Phys Lett 83, 4550–4552 (2003).
9. Menendez, E. et al. Direct magnetic patterning due to the generation of ferromagnetism by selective ion irradiation of paramagnetic FeAl alloys. Small 5, 229–234 (2009).
10. Bali, R. et al. Printing nearly-discrete magnetic patterns using chemical disorder induced ferromagnetism. Nano Lett 14, 435–441 (2014).
11. McCord, J., Schultz, L. & Fassbender, J. Hybrid soft-magnetic lateral exchange spring films prepared by ion irradiation. Adv Mater 20, 2090–2093 (2008).
12. Kim, S. et al. Nanoscale patterning of complex magnetic nanostructures by reduction with low-energy protons. Nature nanotechnology 7, 567–571 (2012).
13. Kim, S., Lee, S. & Hong, J. An array of ferromagnetic nanoislands nondestructively patterned via a local phase transformation by low-energy proton irradiation. ACS Nano 8, 4698–4704 (2014).
14. Fassbender, J. & McCord, J. Magnetic patterning by means of ion irradiation and implantation. J Magn Magn Mater 320, 579–596 (2008).
15. Franken, J. H., Swagten, H. J. & Koopmans, B. Shift registers based on magnetic domain wall ratchets with perpendicular anisotropy. Nature nanotechnology 7, 499–503 (2012).
16. Oby, B. et al. A micro-structured ion-implanted maginonic crystal. Appl Phys Lett 102, 202403 (2013).
17. Sort, J. et al. Periodic arrays of nanoscale and sub-nanoscale magnetic structures prepared by nanoindentation of a nonmagnetic intermetallic compound. Adv Mater 18, 1717–1720 (2006).
18. Zheng, M. et al. Magnetic nanodot arrays produced by direct laser interference lithography. Appl Phys Lett 79, 2606–2608 (2001).
19. Hayashi, M. et al. Spatial control of magnetic anisotropy for current induced dominant domain wall injection in perpendicularly magnetized CoFeB vertical bar MgO nanostructures. Appl Phys Lett 100, 192411 (2011).
20. Li, S. P. et al. Magnetic domain confinement by anisotropy modulation. Phys Rev Lett 88, 087202 (2002).
21. Piramanayagam, S. N. Perpendicular recording media for hard disk drives. J Appl Phys 102, 011301 (2007).
22. Gutleisch, O. et al. Magnetic materials and devices for the 21st century: stronger, lighter, and more energy efficient. Adv Mater 23, 821–842 (2011).
23. Johnson, M. T., Bloemen, P. J. H. & denBroeder FJA, deVries J. J. Magnetic anisotropy in metallic multilayers. Rep Prog Phys 59, 1409–1458 (1996).
24. Wang, K. L., Alzate, J. G. & Amiri, P. K. Non-perturbative spintronic memory: STT-RAM and beyond. J Phys D-Appl Phys 46, 074003 (2013).
25. Thomas, L. et al. Perpendicular spin transfer torque magnetic random access memories with high spin torque efficiency and thermal stability for embedded applications (invited). J Appl Phys 115, 172615 (2014).
26. Pommier, I. et al. Magnetization Reversal in Ultrathin Ferromagnetic Films with Perpendicular Anisotropy - Domain Observations. Phys Rev Lett 65, 2054–2057 (1990).
27. DenBroeder, F. J. A., Kuiper, D. & Draaisma H. J. G. Effects of Annealing and Ion-Implantation on the Magnetic-Properties of Fe/Pd Multilayers Containing Ultrathin Co. IEEE Trans Magn 23, 3696–3698 (1987).
28. Lin, M. T. et al. Giant enhancement of magneto-optical response and increase in perpendicular magnetic anisotropy of ultrathin Co/Pt(111) films upon thermal annealing. J Vac Sci Technol A 17, 3045–3050 (1999).
29. Victoria, R. H., Willoughby, S. D., Maclaren, J. M. & Xue, J. H. Effects of grain boundaries on magnetic properties of recording media. IEEE Trans Magn 39, 719–715 (2003).
30. Kelly, P. J., Ogrady, K., Mayo, P. I. & Chantrell, R. W. Switching Mechanisms in Cobalt-Phosphorus Thin-Films. IEEE Trans Magn 35, 3880–3883 (1989).
31. Wu, X. W., van de Veerdonk, R. J. M., Chantrell, R. W. & Weller, D. Delta M study of perpendicular recording media. J Appl Phys 93, 6760–6762 (2003).
32. Zhang, H. W. et al. Investigation on intergrain exchange coupling of nanocrystalline permanent magnets by Henkel plot. Appl Phys Lett 82, 4098–4100 (2003).
33. Iguchi, M., Hara, M., Nakamura, A. & Sugita, Y. Micromagnetic simulation of magnetic cluster, thermal activation volume, and medium noise in perpendicular recording media. IEEE Trans Magn 39, 1897–1901 (2003).
34. Piramanayagam, S. N. & Srinivasan, K. Recording media research for future hard disk drives. J Magn Magn Mater 321, 485–494 (2009).
35. Skomski, R. Nanomagnetics. J Phys-Condens Matter 15, 8381–R896 (2003).
36. Greaves, S. J., Muraoka, H., Sugita, Y. & Nakamura, Y. Intergranular exchange pinning effects in perpendicular recording media. IEEE Trans Magn 35, 3772–3774 (1999).
37. Zablotskii, V., Ferre, J. & Maziewski, A. Model for domain wall pinning by randomly distributed nanosized defects in ultrathin magnetic films. J Phys D-Appl Phys 42, 125001 (2009).
38. Schuller, I. New Class of Layered Materials. Phys Rev Lett 44, 1597–1600 (1880).
39. Janssen GCAM, Abdalla M. M., van Keulen, F., Pujada, B. R. & van Venrooy, B. Celebrating the 100th anniversary of the Stoney equation for film stress: Developments from polycrystalline steel strips to single crystal silicon wafers. Thin Solid Films 517, 1858–1867 (2009).
40. Thornton, J. A. & Hoffman, D. W. Stress-Related Effects in Thin-Films. Thin Solid Films 171, 51–31 (1989).
41. Vinci, R. P., Zielinski, E. M. & Bravman, J. C. Thermal Strain and Stress in Copper Thin-Films. *Thin Solid Films* **262**, 142–153 (1995).
42. Hyun, S., Vinci, R. P., Fahey, K. P. & Clemens, B. M. Effect of grain structure on the onset of diffusion-controlled stress relaxation in Pt thin films. *Appl Phys Lett* **83**, 2769–2771 (2003).
43. Hazzledine, P. M. & Schneibel, J. H. Theory of Coble Creep for Irregular Grain Structures. *Acta Metall Mater* **41**, 1253–1262 (1993).
44. Weller, D *et al*. Growth, structural, and magnetic properties of high coercivity Co/Pt multilayers. *J Appl Phys* **89**, 7525–7527 (2001).
45. Gan, DW *et al*. Isothermal stress relaxation in electroplated Cu films. I. Mass transport measurements. *J Appl Phys* **97**, 103531 (2005).
46. Chang, C. Y. & Vook, R. W. The Effect of Surface Aluminum-Oxide Films on Thermally-Induced Hillock Formation. *Thin Solid Films* **228**, 205–209 (1993).
47. Roder, H., Schuster, R., Brune, H. & Kern, K. Monolayer-Confined Mixing at the Ag-Pt(111) Interface. *Phys Rev Lett* **71**, 2086–2089 (1993).
48. Jin, Z. Q. & Liu, JP. Rapid thermal processing of magnetic materials. *J Phys D-Appl Phys* **39**, R227–R244 (2006).
49. Yamane, H., Maeno, Y. & Kobayashi, M. Annealing Effects on Multilayered Co/Pt and Co/Pd Sputtering Films. *Appl Phys Lett* **62**, 1562–1564 (1993).
50. Denbroeder, F. J. A., Janssen, E., Mud, A. & Kerkhof, JM. Co/Ni Multilayers with Perpendicular Magnetic-Anisotropy. *J Magn Magn Mater* **126**, 563–568 (1993).
51. Nguyen, V. D. *et al*. Detection of Domain-Wall Position and Magnetization Reversal in Nanostructures Using the Magnon Contribution to the Resistivity. *Phys Rev Lett* **107**, 136605 (2011).

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**Author Contributions**

K.F.H. designed the experiments and fabricated the magnetic patterns; K.F.H., J.W.L. and L.W.W. performed the magnetic analysis; K.F.H., J.W.L., H.H.L. and C.H.L. analyzed the data of magnetic properties and derived the analytical model; K.F.H., J.W.L., L.W.W., Y.C.H. and W.W.C. prepared the sample for structural analysis and performed structural analysis; C.Y.H., J.Y., M.T.C. and S.C.L. performed the Cs-corrected STEM; K.F.H., J.W.L., J.Y. and C.H.L. discussed and proposed the kinetics model; K.F.H., H.H.L., J.Y., J.W.L. and C.H.L. discussed and wrote the manuscript together. C.H.L. oversaw the project including analysis, discussion, and conceptual guidance; all authors discussed the results and commented on the manuscript.

**Additional Information**

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