Investigating the Super Paramagnetic Limits of Ferromagnetic Nanowires

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ABSTRACT----Highly sensitive focused magneto optical Kerr effect magnetometer (MOKE) has been used to explore the nanoscale limit of state of the art electron beam lithography (EBL) patterned permalloy (NiFe) nanowires. The nanowires are 20 µm long arranged in 20 µm × 20 µm arrays. Permalloy of different thicknesses, which includes 10, 8, 6, 4 and 3.5 nm were deposited on 5 different samples containing these arrays. Results from 10 nm thick permalloy nanowires shows superparamagnetic transition at 85 ± 5 nm wire width. For 6 nm thick array, superparamagnetic transition occurs at 125 ± 25 nm. We therefore show that with 10 nm thickness, instead of 6 nm thickness, more miniaturisation of nanowire width can be achieved before the onset of superparamagnetism. This may be seen as very efficient in achieving increased storage density in memory devices.

Keywords---Permalloy, nanowires, coercivity, superparamagnetic transition

1. INTRODUCTION

The inherent property of ferromagnets to align their magnetic moment strongly in the direction of applied magnetic field is threatened as the magnets are miniaturized down to sub hundreds and tenths of nanometer. Apart from their strong magnetization in the non miniaturized state, they retain internally induced magnetic field even when the previously induced external field is removed [1]. Paramagnets on the opposite has randomly arranged and weakly coupled magnetic moments and do not retain their internal induced magnetic field once the applied field is removed [2]. Superparamagnetism occurs in further miniaturized ferromagnets [3] - a phenomenon whereby ferromagnetic materials respond to applied external magnetic field in a similar way to that of a paramagnet.

Ferromagnetic nanowires have received more attention due to their applications in magnetic logic devices and racetrack memory. They are also used to achieve increased storage density in memory devices and long term stability of memory or recording. As the width of ferromagnetic nanowires decreases, intensity of the magnetic field needed to reduce their magnetisation to zero after saturation magnetisation has been achieved (coercivity) increases rapidly. In other words, ferromagnetic nanowire experiences an inverse width relationship as show in equation 1 with this magnetization reversal from film thicknesses ≥ 3.5 nm [3]. This shows the tremendous effect of geometry of a magnetic sample on the magnetization reversal.

\[ H_c = a + \frac{(3M_s d)}{W} \]  

(1)

where W is the width of the nanowire, M_s is saturation magnetization, H_c is coercive field and d is thickness of the magnetic material deposited on the wire. The inverse width dependence occurs because magnetization reversal is controlled by a small nucleus that spreads across the width of a nanowire but does not depend on the length of the wire. However, with further reduction in the width and thickness of a nanowire, size of the controlling nucleus approaches superparamagnetic limit, which is a region of sudden decrease in coercivity with decreasing wire width. As a result, the inverse width relationship with magnetization reversal breaks down. A curve equivalent to that shown in Figure 1 is an example of superparamagnetism (a sudden decrease in coercivity with decreasing wire width) for 2.5 nm NiFe lines [3].
Figure 1: Trend showing sudden decrease in coercivity with decreasing wire width for a very thin wire of 2.5 nm superparamagnetic transition [3].

2. EXPERIMENT

State of the art electron beam lithography (EBL) can be used to make nanowire patterns with diameter down to tens of nm. In this research, room temperature EBL patterned 20 µm × 20 µm arrays of 20 µm long Permalloy (NiFe) nanowires has been fabricated. The width of the nanowires ranges between 10 to 500 nm, which includes 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 150, 200, 300, 400 and 500 nm with varying thickness of 3.5 to 10 nm and wires of the same width arranged in the same array. Nanowires in an array are separated by twice their width while 50 µm gaps separate one array from the other.

The whole process fabrication of the NiFe nanowires involved nanowire design with LibreCAD, patterning with electron beam lithography (EBL), deposition of permalloy with thermal evaporator, measurement with magneto optical Kerr effect (MOKE) magnetometer and imaging and section analysis with atomic force microscope (AFM). A concise and comprehensive list of the materials and equipment used for each stage of the research is given in Table 1 below.

<table>
<thead>
<tr>
<th>S/N</th>
<th>ACTIVITY</th>
<th>METHOD/ EQUIPMENT USED</th>
<th>CHEMICAL USED</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Design of nanowires</td>
<td>Computer aided design-LibreCAD/ Snap to the grid command.</td>
<td>Not applicable</td>
</tr>
<tr>
<td>2</td>
<td>Patterning of the nanowires</td>
<td>Electron beam lithography (EBL), silicon substrate, spin coater, hot plate for soft baking.</td>
<td>Acetone for cleaning of silicon substrate, isopropyl alcohol (IPA) for washing off of acetone, Polymethylmethacrylate (PMMA) as the positive resist, silicon substrate.</td>
</tr>
<tr>
<td>3</td>
<td>Deposition of permalloy (NiFe)</td>
<td>Thermal evaporator</td>
<td>Polymethylmethacrylate (PMMA) as a mask and for sticking the silicon substrate onto the sample holder, NiFe as the magnetic material deposited on the patterned silicon substrate, acetone for lift-off, isopropyl alcohol (IPA) for rinsing of the acetone, hand gloves and tweezers.</td>
</tr>
<tr>
<td>4</td>
<td>Measurement of magnetic properties</td>
<td>Focused magneto optical Kerr effect (Focused MOKE) magnetometer</td>
<td>Not applicable</td>
</tr>
<tr>
<td>5</td>
<td>Imaging and section analysis</td>
<td>Atomic force microscope (AFM)/contact mode probe</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

Table 1: List of equipment and materials used in the research

Design of the Magnetic Nanowire Structures

LibreCAD is 2D and geometry computer aided design (CAD) which is able to file in the industrially accepted GDSII format. Snap to grid command contained in it makes it easy for basic shapes to be drawn with high dimension accuracy. With LibreCAD, 20 µm × 20 µm arrays of 20 µm long wires were designed. The wire width ranges from 10 to 500 nm and includes 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 150, 200, 300, 400 and 500 nm with wires of the same width arranged in an array and separated by twice their width. 50 µm gaps separate one array from the other. Different shapes of marker
pad were designed 300 µm away above each array for easy identification of the nanowire arrays during measurement. Figure 2 shows the schematic diagram of the arrangement of the nanowire structures.

Figure 2: Schematic diagram of (a) 20 µm × 20 µm arrays of 20 µm long, *W represents different nanowire width, which includes 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 150, 200, 300, 400, and 500 nm. nanowires of the same width are arranged in the same array.

Patterning the Nanowires
EBL patterning of the nanostructures were performed with Joel JBX 6300FS electron beam lithography system at the University of Leeds, United Kingdom. Its minimum feature size is < 8 nm with beam-electron overlay accuracy of < 25 nm. The accelerating voltage is 100 KV and beam currents from 30 pA to 20 nA. Due to nano dimensions of the designed structures, 5 atomically flat SiO₂ substrates unto which the patterns were transferred were first cleaned in 50ºC acetone for 5 minutes, after which it was rinsed in cold acetone then isopropyl alcohol (IPA) and dried with gaseous nitrogen. 2% solution of PMMA in anisole was spin coated on the substrates at 1000 rpm for 10s. Then the substrates were soft baked at 50ºC for 1 hour. Afterwards, the substrates were transferred into the vacuum chamber of the EBL system and exposed to the electron beam following the computer generated pattern. The exposed substrates were developed by immersion in methyl-isobutyl-ketone (MIBK) in the ratio 1:3 with IPA. Afterwards, they went through ultra sound agitation for 30 seconds.

Permalloy Deposition
Thermal evaporation and sputtering are methods that can be used to deposit magnetic materials unto a substrate. With thermal evaporation, very thin films (mostly metals) can be deposited onto a substrate in a high vacuum environment of about 10⁻⁷ mbar, which allows vapour particles from the target material to travel directly to the substrate. It deposits in the thickness range of angstroms to microns and requires low voltage and high current to function. In sputtering, atoms are removed from a target by impact from ions of inert gas plasma like Argon. A high vacuum chamber is filled with the inert gas and voltage applied across the gas creates plasma. Positive ions generated accelerates to the cathode where material to be deposited is placed and sputters atoms from it, which condenses unto the substrate.

In this research, thermal evaporator was used to deposit permalloy (NiFe) onto 5 different EBL patterned substrates- 1,2,3,4 and 5. Each substrate differs from the other only by the thickness of permalloy deposited on them. Substrates 1,2,3,4 and 5 contains 10, 8, 6, 4 and .5 nm thick NiFe respectively. The thermal evaporator uses tungsten basket. The base pressure of the vacuum chamber was 3 × 10⁻⁷ mbar, which increased to 7 × 10⁻⁶ mbar during evaporation. Lift-off was done for 60 seconds in acetone by ultrasonic vibration and the samples were rinsed in IPA.

Magneto Optical Kerr Effect Magnetometer
Magneto optical Kerr effects in magnetic materials arise due to the optical anisotropy of such materials. These materials have magnetisation within surface domains, which can be influenced by magnetic field [4]. MOKE is similar to Michael Faraday effect where the polarisation of light is rotated through a transparent material subjected to a magnetic field. The operational principle is that when polarised light is incident on a magnetic material, it reflects and changes its polarisation. Change in polarisation is proportional to magnetisation. Therefore, through measuring change in polarisation, magnetisation of the magnetic material of interest can be deduced.
MOKE is a non-destructive technique for measuring magnetization in magnetic materials. Due to its high spatial resolution, which is still developing, and high sensitivity, it remains at the forefront of magnetic nanostructure research. With it, domain wall propagation in ultrathin magnetic nanowires has been imaged [5] and the magnetisation response of the individual structures studied on ultra-fast time scale with a spatial resolution <200 nm [6]. Also, it has been used to study isolated magnetic microstructures [7, 8], behaviour of nanodot chains [9, 10] and even the behaviour of selected regions of separate nanowire structures [11, 12].

For sample imaging, white light that is brought collinear with the laser optical axis by means beam splitter between the objective lens and polariser forms the imaging optical path. Dichroic mirror placed before the CCD chip filters intense laser beam. While the sample surface is imaged unto a CCD camera through the beam splitter placed just before the λ/4 wave plate. Iterative rotation of the analyser first then λ/4 wave until a minimum voltage is achieved, followed by rotation of the analyser by 20º for further voltage reduction sets up the polarisation optics for measurement. Diode-pumped solid state laser (coherent Verdi) with operating wavelength of 532 nm and continuously variable output power ~ 5 W kicks off the measurement procedure. The ring cavity has excellent low stability characteristic of about 0.02 %, measured between 10Hz to 1GHz. Therefore, high power output, half- wave plate /polariser system and neutral density (ND) filter are used in attenuating the laser to obtain stability characteristic close to that of the laser light. The beam polarisation ratio is improved by allowing it pass through beam expander and a high quality Glan-Taylor polarizing prism with extinction ratio of < 10⁻³. The beam focuses on ~ 5 μm diameter spot of the sample. Reflection from the spot is collected with identical chromate doublet lenses with infinite conjugate ratios and 500 mm focal length. The focused laser is supposed to have perfect or at least near perfect Gaussian profile in the plane of incidence but does not because the sample is mounted at 45º to the optical axis. This is achieved by having the objective lens adjusted until the focused spot has high central intensity that is sufficient without significant higher order mode being noticed. The laser beam is then directed to a Glan-Taylor polarising analyser through a λ/4 wave guide retardation plate, which introduces a relative phase shift of 90º in the reflected light. Finally, Silicon photodiode detects the reflected beam, providing Kerr signal.

Photodiodes are connected to the lock in amplifier, which features ±2V offset adjustment. The offset is adjusted as the detectors are illuminated until each amplifier gives a net output signal close to zero, though the offset is later added to the signal as part of data acquisition. This improves the effective dynamic range of the measurement. Data averaging occurs in the digital storage oscilloscope before being downloaded to the PC for analysis. Magnetic field is generated in the plane of the sample by electromagnets and iron-core driven by KEPCO power amplifier [13]. TTI function generator defines the waveforms. XYθ stage is a computer controlled translation stage with 55nm lateral resolution where magnetic samples are mounted. CCD camera image guides in placing the sample well. The sample is raster scanned through a two-dimensional grid beneath the laser spot, which is recorded with the signal from the lock-in amplifier for each position in the grid. The signal allows susceptibility map of the sample to be generated. Using this, a specific area in the magnetic structure can be studied as well as individual magnetic nanostructures identified [14, 15].

3. RESULTS

To check for superparamagnetic transition (SPT) in the nanowires, plots of inverse of the designed nanowire widths versus coercivity H, for samples 1 (10 nm thick) and 3 (6 nm thick) wires were made as shown in Figure 3. Following the trend of inverse width relationship with coercivity discussed earlier, SPT is evident in the plots. 10 nm thick nanowires in Figure 3 (a) shows SPT between 0.0111 and 0.0125 nm⁻¹. This gives the wire width of about 85 ± 5 nm. The SPT of 85 ± 5 nm can also be clearly seen in the graph of Hc (Oe) versus width (nm) shown in figure 3 (b). Result from 6 nm thick nanowires is shown in figure 4. Figure 4 (a) shows SPT between 0.007 and 0.01 nm⁻¹, which is about 125 ± 25 nm. The SPT of 125 ± 25 nm can also be seen clearly in the graph of Hc (Oe) versus width (nm) shown in figure 4 (b).
Figure 3: Result from sample 1 (10 nm thick nanowires). (a): Plot of coercivity, $H_c$ (Oe) versus inverse of nanowire width showing superparamagnetic transition between 0.0111 and 0.0125 nm$^{-1}$ (b) Plot of coercivity, $H_c$ (Oe) versus nanowire width showing SPT at 85 ± 5 nm.
Figure 4: Result from sample 2 (6 nm thick nanowires). (a): Plot of coercivity, $H_c$ (Oe) versus inverse of nanowire width showing superparamagnetic transition between 0.007 and 0.01 nm$^{-1}$ (b) Plot of coercivity, $H_c$ (Oe) versus nanowire width showing SPT at 125 ± 25 nm.

4. CONCLUSION
We have investigated superparamagnetic transition in permalloy nanowires made by Electron Beam Lithography and Thermal Evaporation. We show that superparamagnetism sets in at wire width of 85 ± 5 nm and 10 nm thickness. Also, for 6 nm thickness, superparamagnetic transition sets in at 125 ± 25 nm wire width. We therefore show that with 10 nm thickness, instead of 6 nm thickness, more miniaturisation of nanowire width can be achieved before the onset of superparamagnetism. This may be seen as very efficient in achieving increased storage density in memory devices.

5. ACKNOWLEDGEMENTS
We thank Tertiary Education Trust Fund (TETFund) for sponsoring this work.
6. REFERENCES