STRUCTURAL AND MAGNETIC PROPERTIES OF FE-OXIDE LAYERS PREPARED BY INKJET PRINTING ON SI-SUBSTRATE

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Abstract

Magnetic films with two, four, six, and eight layers were prepared using 2D inkjet printing on the 333 K heated Si-substrates at ambient conditions. The microstructure, phase composition of layers, and magnetic studies were investigated using wide spectrum of experimental methods with the aim to find the optimal conditions for producing magnetic sensors by digital printing technology in the near future. As the magnetic compound, the hematite powder was chosen and its milling with Si-binder dissolved in dowanol using glass balls and vial to prevent contamination was done to prepared suspension appropriate for printing. It is shown that hematite transformed markedly into maghemite, thus the layers consisted of small amount hematite and dominant maghemite phases determining the magnetic properties. From the viewpoint of magnetic behavior, the highest saturation magnetization and the lowest coercivity was obtained at the sample with 8 layers.

Keywords: Inkjet printing, hematite, physical properties, testing methods

1. INTRODUCTION

The reduction of materials scale leads to profound changes in their inner and surface microstructure and accompanying modification of their physical properties, e.g., electronic, optical, magnetic. In the case of magnetic nanoparticles, the miniaturization often means a deviation from the established laws governing the magnetic phenomena observed in bulk magnetic materials. The current research is oriented to preparation of magnetic nanoparticles using various technological procedures, to their microstructure at the nanometric scale, and to correlations between the microscopic and macroscopic properties. The usual and/or innovative experimental methods allow determination of anomalous or enhanced properties as giant magnetoresistance, coercivity, remanence and magnetocaloric effect enhancements and quantum tunneling of the magnetization, etc. This gives rise to enormous technological usage in, e.g., next generation of very high-density magnetic information storage devices, magnetic sensors, permanent magnets, biocompatible products, etc.

Development of 2D inkjet printing technology enabled to deposit thin layers of materials onto rigid and flexible surfaces. It presently attracts great interest predominantly in the prohexanolduction of low-cost and large-area electronic devices [1]. On the other hand, its usage in the production of magnetic devices is at the very beginning. Among pioneering works, the studies of the magnetic film prepared by a digital printing have to be mentioned. The magnetic film was prepared from the magnetite nanoparticles covered with oleic acid and investigated with the aim to realize magnetic cores for Radio Frequency Identification Resonators [2].

Present work concentrates on the possibility for the production of thin magnetic layer on Si-plate using 2D inkjet printing technology at the temperature slightly above room temperature under ambient atmosphere. The main aim is to produce films of different thickness and to characterize their structure and magnetic properties,
and to predict the optimal conditions of production in relation to physical properties for future application in magnetic sensors.

2. EXPERIMENTAL DETAILS

As an input magnetic material, the commercial hematite nanoparticles of dimension < 50 nm (Sigma-Aldrich, Steinheim, Germany) was used. The hematite nanoparticles dispersed in dowanol were initially milled with polysiloxane binder dissolved in absolute ethanol [3] using balls in vial; both made of glass to prevent contamination. The aim was to decrease the particle size and to obtain the acceptable suspension. Prior printing on the 333 K heated Si-substrate at ambient conditions the suspension was in addition diluted with hexanol. The Si-wafers of different size, corresponding to demands of the subsequently applied experimental methods, were prepared by cutting Si-wafers (100) using spark erosion. The smallest dimension 2.5 mm x 2.5 mm was used for magnetic measurements, the larger ones, 10 mm x 10 mm, for scanning electron microscopic observations (SEM), X-ray diffraction (XRD), and Mössbauer (MS) measurements.

2D inkjet printing equipment Dimatix Materials Printer DMP-2850 was used for the formation of iron-oxide layers of different thickness; two, four, six, and eight layers (L) were chosen and in conformity with it the samples were denoted 2L, 4L, 6L, and 8L. The approximate thickness of layers was determined using profilometer Dektak XT, Bruker.

The morphology of samples was followed by scanning electron microscope MIRA3 (Tescan, Brno) at accelerating voltage of 30 kV.

X-ray diffraction patterns were measured at room temperature (RT) using an EMPYREAN diffractometer (Panalytical) with Co Kα radiation (λ = 0.17902 nm) in the 2θ range from 15° to 130° in steps of 0.026° and time per step ≈400 s. The evaluation of powder patterns was realized by the Rietveld structure refinement method [4] by semi-automatic mode using the HighScore Plus program and the ICSD database of inorganic and related structures [5]. The phase composition and the lattice parameters were obtained from the pattern analysis.

Fe Mössbauer spectra at room temperature were taken in the transmission geometry using γ-rays and scattering geometry by detecting 7.3 keV conversion electrons (CEMS, [6]) with penetration depth of about 200 nm. Obtained spectra were analyzed using CONFIT program package [7] and calibration of velocity scale was done by a standard thin α-iron foil at RT. The transmission geometry was used for measurement of the hematite-suspension soaked in cellulose. Scattering geometry was applied for measurements of the final 2L and 8L samples.

The magnetic measurements were performed using a Physical Property Measurement System (PPMS) Quantum design, Inc. The hysteresis curves were taken at 300 K and 2 K in magnetic field of ± 800 kA/m. The zero field-cooled (ZFC) and field-cooled (FC) curves were measured between 2 K and 300 K in magnetic field of 8 kA/m.

3. RESULTS AND DISCUSSION

3.1. Sample characterization

The scanning electron microscope images from all studied samples appear very similarly without any dependence on the layer thickness, as shown in Figure 1 representing the 8L (a) and 2L (b) samples. The thickness obtained from profilometry scans (not shown here) linearly increased from approximately 2.2 μm for the 2L sample to 7.5 μm for the 8L sample. The surfaces of all samples yield small agglomerated nanoparticles of size below 50 nm. The top layers are not fully homogeneous and, in some spots, the pores of dimensions between 50 nm and 300 nm can be observed. While in a case of the 8L sample the sub-layers are visible, the similar small pores observed at the 2L sample often detect the Si-substrate.
The X-ray diffraction measurements of all samples resulted in very similar patterns and therefore only those for the 2L and 8L are depicted in Figure 1 (c) as representatives. The diffractograms analysis resulted in a presence of dominant magnetite/maghemite and only minor content of hematite. This stimulated the Mössbauer measurements of the hematite suspension using transmission geometry and of the 2L and 8L samples using scattering geometry. The spectra are depicted in Figure 1 (d) and the results followed from experimental data analysis of both methods are summarized in Tables 1 and 2, respectively.

Table 1 Phase analysis of X-ray diffractograms; $A$ – phase content, $a, b, c$ – lattice parameters

|     | Phase | $a$, $b$, $c$ (nm) | $A$ (wt%) | $a$ (nm) |
|-----|-------|-------------------|-----------|----------|
| 2L  |       | 0.5033(6), 1.3689(20) | 97.5      | 0.8346(6) |
| 4L  |       | 0.5037(9), 1.3926(38) | 94.3      | 0.8336(4) |
| 6L  |       | 0.5038(6), 1.3669(22) | 94.0      | 0.8336(6) |
| 8L  |       | 0.5036(6), 1.3709(23) | 95.6      | 0.8350(9) |

Hematite is a common iron oxide with a formula of $\alpha$-Fe$_2$O$_3$. It was supposed to be the initial iron-oxide for the present studies; however, the XRD measurements have shown dominant maghemite phase. It is well known that hematite is highly sensitive to various effects; e.g., pressure, external magnetic field, structural defects, impurities, and grain size, influencing its physical properties. For its determination in measured diffractograms the ICSD 15840 sheet with input values of the lattice parameter, $a = b = 0.5038$ nm and $c = 1.3772$ nm, was used. As it follows from Table 1 its content in all samples is low, which reflects its partial transformation into
dominant magnetite/maghemite phase. It could be very probably caused by induced stresses during milling in a process of the iron-oxide suspension preparation. The magnetite, Fe₃O₄, is crystallographically isomorphous with maghemite, γ-Fe₂O₃, and both these iron-oxides have cubic structure of similar lattice parameters. Therefore, their determination from diffractogram is not straightforward. For the Rietveld analysis the ICSD data sheets 26410 for magnetite (a = 0.83941 nm) and 172905 for maghemite (a = 0.84058 nm) were used. The resulted values of lattice parameters of both phases did not allow to distinguish between magnetite and maghemite unambiguously. Therefore, the values in Table 1 correspond to weighted mean average.

Table 2 Mössbauer characteristics deduced from the transmission spectrum of the input iron-oxide-suspension and conversion scattering spectra of the 2L and 8L samples; A – phase content, B – hyperfine induction, δ – isomer shift, and Δ – quadrupole shift

| Sample                  | hematite   | maghemite | superparamagnetic |
|-------------------------|------------|-----------|-------------------|
|                         | A (%)      | B (T)     | δ (mm/s)          | Δ (mm/s) | A (%) | δ (mm/s) | Δ (mm/s) |
| Iron-oxide              | 10.3       | 50.5      | 0.359             | -0.183   | 75.9  | 49.1     | 0.257    | 0.027    | 13.8  | 0.179    | 0.553    |
| suspension              |            |           |                   |          |       |          |          |          |       |          |          |
| 2L                      | 7.6        | 51.5      | 0.396             | -0.143   | 87.3  | 43.9     | 0.335    | 0.008    | 5.1   | 0.384    | 0.813    |
| 8L                      | 10.1       | 51.3      | 0.412             | -0.164   | 79.8  | 48.1     | 0.331    | 0.011    | 10.1  | 0.372    | 1.162    |

57Fe Mössbauer spectra recorded at RT were analysed using a conventional least-square refinement method with combination of sextets and doublets of Lorentzian lines. The results shown in Table 2 reflect about 10 % of hematite in the input iron-oxide-suspension represented by sextet of hyperfine parameters, isomer shift and quadrupole splitting, in good agreement with Ref. [9]. The rest is ascribed to magnetically ordered maghemite which documents hematite transformation owing to milling. This confirms the XRD observations and it is supported by Randrianantoandro and co-workers [10] reporting direct phase transformation from hematite to maghemite during high energy ball milling. The doublets in the middle of spectrum can be ascribed to superparamagnetic (SP) portion of iron-oxides in a form of very small nano-grains (<10 nm). Slightly lower content of this component was detected also in scattering spectra taken from the 2L and 8L samples in which dominant component was again maghemite. The values of hyperfine induction of hematite correspond to literature but those of maghemite, mainly at the 2L sample, are lower. It could be due to slight distortion of crystallographic sites mainly at the particle surfaces and insufficient energy needed for the lattice formation and vacancies reordering. For more detailed analysis, the Mössbauer measurements at low-temperature and, eventually, in an external magnetic field should be helpful.

3.2. Magnetic results

The nearly square hysteresis loops of all samples measured at room and low temperatures are shown in Figure 2. The changes of magnetic characteristic, saturation and remnant mass magnetizations $M_s$ and $M_r$ and coercivity $H_c$, in dependence on number of layers are presented in Figure 3. The mean values of the coercivity $H_c$ and remnant magnetization $M_r$ were obtained from the polynomial fitting of hysteresis loops between -60 and 60 kA/m and $M_s$ was calculated from both branches in saturation (above ≈ 700 kA/m). The experimental points were fit by linear functions (dashed lines) better to see tendency of parameter changes. The dependences (Figure 3) reveal that increasing number of layers (film thickness) leads to an increase of the saturation and remnant magnetizations at both measuring temperatures while the $H_c$ embodies an opposite tendency. Such behaviour is not general. For example, in Permalloy film study is the behaviour of magnetic characteristics different bellow and above estimated critical thickness [8]. For the 8L sample the saturation mass magnetization is slightly lower compared to values (60-80 Am²/kg) presented for maghemite in literature (e.g. [9]) probably due to influence of the low RT saturation magnetization of weakly ferromagnetic hematite (≈ 0.3 Am²/kg, [9]) and reasons mentioned above in discussion of Mössbauer results.
Figure 2 Hysteresis loops of 2L, 4L, 6L and 8L samples measured at 300 K (a) and 2 K (b); ZFC-FC curves (c)

The $H_c$ and $M_r$ dependences on number of layers are not so unique compared to $M_s$ due to their high structural sensitivity. The ZFC and FC curves for all samples in Figure 3c are similar. In agreement with XRD and MS observations they do not provide any sign of Morin temperature being for hematite 260 K or lower in dependence on particle size.

Figure 3 Dependences of saturation magnetization (a), remnant magnetization (b) and coercivity (c) on number of layers taken from hysteresis loops measured at 2 K and 300 K

4. CONCLUSION

The present investigations were devoted to application of the 2D inkjet printing procedure in preparation of magnetic film on the Si-substrate. The hematite powder dispersed in dowanol and mixed with siloxane binder dissolved in absolute ethanol was milled using glass balls and vial with the aim to form suspension applicable for digital printing. The samples with two, four, six, and eight laeyrs of magnetic film were produced and investigated with numerous experimental methods. The scanning electron microscope observations yielded very similar microstructure formed by particles of size below 50 nm and their agglomerations among them sometimes the pores were visible. These pores detected the Si substrate at the two-layers sample while at the eight-layers sample the bellow laying particles were seen in the pores. It means that cross-sectional homogeneity was not quite satisfactory. The analysis of the X-ray diffraction patterns supporred by Mössbauer phase analysis has found dominant magnetite/maghemite phase and only small amount of the initial hematite in layers of all sample. This transformation of hematite into magnetite/maghemite was found to be initiated during milling in procedure of the suspension preparation. This was subsequently confirmed by high values of saturation magnetization and the measurements of the ZFC-FC curves at which no Morin transition, characteristic for hematite, was observed. From the magnetic viewpoint, the highest saturation magnetization and the lowest coercivity was at the sample with 8 layers.
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