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Impact of aligned carbon nanotubes array on the magnetostatic isolation of closely packed ferromagnetic nanoparticles

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a b s t r a c t

We investigate the influence of carbon nanotubes (CNT) aligned array on the magnetic properties of ensemble of densely packed Co nanoparticles (NPs) embedded inside CNT. Each CNT contains only one nanosized Co. Such a special structure was formed by catalyst chemical vapor deposition (CCVD) activated by current discharge plasma and hot filament. The Co NPs, previously deposited onto SiO₂/Si substrate, acted as a catalyst. By varying the parameters of the CCVD process, we were able to also sputter the substrate instead of CNT growth. Co NPs were used as a mask and the structure of Si-based nanocones with Co NPs on the top of each cone was formed. Exhaustive investigation of the structural, morphology and crystalline properties of Co nanoparticles were performed. The magnetic properties of two kinds of samples, Co on the Si-based nanocone and Co inside CNT, were differ drastically. In the former case, the magnetic anisotropy of thin-film-type has been observed with large magnetic domains. Whereas for the Co-CNT samples ferromagnetic NPs were magnetically isolated. It was established that the magnetic anisotropy of nanosized Co plays more dominant role than the dipole interaction between Co NPs. The role of the CNT container in this is discussed.

1. Introduction

At present, it has been unequivocally established that the properties of magnetic nanoparticles (NPs) differ significantly from bulk phase. For example, when the ratio of surface spins to the total number of NP's spins increases, surface effects dominate the overall NP properties [1]. Considering an ensemble of NPs it is worth underlying the interplay between interparticle direct or indirect exchange coupling and magnetic anisotropy, magnetic anisotropy and dipole-dipole interparticle interaction, intraparticle exchange coupling between surface and core. In other words, the interplay between collective effects and internal NP features determines the macroscopic properties of the whole system. In particular, in aggregated magnetic NPs, when they are in close contact, exchange

interactions could determine their characteristics. On the other hand, strong dipole-dipole interaction (DDI) between closely packed ferromagnetic NPs leads to collective effects [2e4]. However, for many applications, an ensemble of noninteracting magnetic NPs is required.

There are numerous approaches to create an ensemble of isolated (non-interacting) magnetic NPs. A variety of parameters which must be controlled, like NP parameters (chemical nature, distribution, mean size, concentration, interparticle spacing, shape, agglomeration), matrix material parameters (chemical nature, conductivity, permeability, method of NPs embedding into the matrix) create a huge space for experimental research in this interdisciplinary field [2,5e15]. The general tendency is that the increase of the interparticle spacing reduces the interparticle dipolar coupling and ultimately creates a system of non-interacting NPs [16,17]. A serious problem arising in the synthesis process however is agglomeration of nanoparticles. Various chemical methods of their fabrication, such as co-precipitation, thermal decomposition, hydrothermal, sol-gel and other approaches [13,18] can only change the effectiveness of the agglomeration but cannot avoid it completely. Moreover, the agglomeration prevents the

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fabrication of an ensemble of closely packed magnetically isolated NPs, which is a great challenge for modern magnetoelectronics and spintronics.

Recently it has been demonstrated that carbon nanotubes (CNT) decorated by ferromagnetic nanoparticles can provide a very useful platform for studying various types of interparticle interaction, as well as impact of the CNT matrix on the nanoparticle's arrangement and interplay between individual and collective properties [6,19]. Indeed, when one produces oriented CNT arrays by chemical vapor deposition using 3d metals (Fe, Ni, Co) as a catalyst (CCVD), such samples can be considered as a magnetic nanocomposite with ferromagnetic NPs embedded inside CNT matrix. By floating CCVD, depending on the concentration of the catalyst, one can get ferromagnetic NPs located only inside the inner CNT channels for low concentration, or both distributed inside and outside the CNT walls for high catalyst concentration [19,20]. In the latter case NPs are strongly coupled via direct and indirect exchange interaction [21e23], whereas in the former case there are indications of magnetically isolated NPs [24]. This method creates nanotubes with many ferromagnetic NPs inside the inner channel of the same nanotube and the mean distance between adjacent nanoparticles is order of hundreds of nanometers [23]. Nevertheless, the characteristic feature of the exchange coupling between these NPs as observed [21e23,25]. It was suggested that due to high spin orbit interaction in defective CNTs the indirect exchange coupling via CNT inner shell is the reason for this [23,26].

It is natural to expect that if each CNT contains only one ferromagnetic nanoparticle inside an inner channel and all CNTs are perfectly aligned, but are not in contact with each other, then one can get an ensemble of densely packed ferromagnetic NPs without the possibility of any exchange coupling between them. Such kind of samples can be synthesized by the plasma-enhanced CCVD (PE-CCVD) process on flat substrate onto which NPs of a 3d metal were preliminary deposited. The PE-CCVD growth is activated by a direct current discharge plasma (PE CCVD) or by heating by hot filament (HF CCVD), or by both processes (PE HF CCVD). As it was demonstrated in Refs. [27,28], application of both modes of gas activation is the best adapted to grow vertically aligned carbon nanotubes using Co NPs as a catalyst. However, with a high carbon concentration in the PE HF CCVD mode, competition between carbon deposition and substrate sputtering may occur [27]. Under appropriate conditions, it is then possible to grow new highly oriented materials with a single individual ferromagnetic Co NP on top.

In this paper, we consider nanomaterials synthesized by PE HF CCVD on SiQ/Si substrates covered with Co NPs at different carbon concentration during the growth process. We get quite different configurations of Co NPs and we investigate their magnetic properties. After this Introduction, the paper is organized in 3 sections: the experimental preparation procedure and sample characterization are described in Section 2. We underline especially that, depending on the parameters of the PE HF CCVD, it is possible to obtain a single Co nanoparticle embedded on top of a silicon oxynitride (SiN_xO_y) nanocone and a single Co quasi-nanocylinder inserted inside CNTs. By means of electron diffraction we prove the existence of both fcc and hcp Co. The competition between intrinsic and long-range DDI is experimentally investigated in Section 3. For that we start from magnetic force microscopy (MFM) measurements, afterwards we analyze field cooled and zero field cooled (FC-ZFC) temperature dependence of the magnetization. From both results we unambiguously obtain that for Co-CNT samples inclusions are non-interacting while in the Co-SiN_xO_y case the DDI between Co nanoparticles dominates. Then the magnetization M(H) loops are investigated. To minimize the effect of temperature, data were acquired at T $\frac{1}{4}$ 2 K. We show that for Co-SiO_xN_y samples the in-plane magnetic anisotropy dominates, which confirms

strong DDI within the plane. Reversely for Co-CNT samples, it is not possible to distinguish the easy axis of magnetization. In Section 4 both magnetic systems are studied and the micromagnetic parameters including the effective magnetic anisotropy constant of Co NPs are estimated within the random anisotropy model (RAM). We show that the field of the random anisotropy exceeds the exchange field in both cases. But the nature of the magnetic anisotropy contribution is still unclear. To solve this problem, we consider the most significant contributions to the obtained effective magnetic anisotropy constant of Co NPs. For that we perform a comparative analysis of the DDI and magnetic anisotropy and we deduce the contribution of the DDI, magnetocrystalline, magnetoelastic and shape anisotropies (MCA, MEA and SA, respectively). We underline that, the contribution of the MEA depends essentially on the type of the crystalline lattice. We separately estimate the contribution of fcc and hcp Co to the elastic stresses. In the latter case different orientations of the hexagonal axis with respect to the CNT alignment are considered. It is obtained that maximum elastic stresses occur when the hexagonal axis is oriented perpendicular to the CNT axis. Even within these conditions Co remains elastic and the corresponding deformations of the CNT lattice do not exceed 2%. At the end of this section, we show the results of the micromagnetic simulations which support the idea about the perpendicular orientation of the hexagonal axis of hcp Co. The article ends with some concluding remarks and outlook.

2. Experimental and Co NP's characterizations

A SiO₂ layer of 5 nm thickness was first deposited onto Si(100) substrate by an electron cyclotron resonance plasma process. The two samples preparation are then processed differently.

2.1. Nanoparticles on top of nanocones: synthesis and characterization

An ultrathin Co film (2.5 nm) was sputtered by DC magnetron. The buffer SiO₂ layer prevents the formation of Co silicide. In addition, there is large difference in surface energies of SiO₂ and Co which facilitates the coalescence of former Co atoms with the formation of small nanoparticles. Then the substrate was transferred into an UHV CVD reactor and softly elaborated by slow thermal reaction up to 973 K with a heat rate of 10 K/min under UHV and above 673 K under a hydrogen/ammonia (80:20) gas flow at 15 mbar. Finally, the metal reduction and the formation of Co NPs were performed in a thermally and plasma-activated mixture of hydrogen and ammonia at 973K during 15 min. As a result, an ensemble of Co NPs was synthesized on the surface of SiO₂/Si substrates. Then, the process of PE HF CCVD was performed in the mixture of C₂H₂:NH₃:H₂ $\frac{1}{4}$ 20:16:64 with a flow rate D_{flow} $\frac{1}{4}$ 100 sccm at 15 mbar. The conditions are summarized in Table 1 for Co catalyst preparation and in Table 2 for nanostructures preparation. As a result, in the presence of heavy nitrogen-based ion, sputtering of the SiO₂/Si substrates readily occurs. Therefore, the resulting sample consists of silicon oxynitride nanocones with the Co NPs merging on the top of each nanocone. However, the Co NPs behaved like a mask during the sputtering process, protecting the area below from sputtering. Also, the sputtering by directional ions allowed the definition of geometrical nanocones centered around the Co NP mask. A SEM image (XL30S-FEG PHILIPPS working at 3 KV) of a typical sample is shown in Fig. 1a and a TEM image (TOPCON 002B working at 200 kV with dry surface scratching) is displayed in Fig. 1b.

Other synthesis conditions (experimental set-up, heating and cooling steps, plasma activation parameters) and Co NPs properties (size histogram and SEM image in top view) are displayed in the

Table 1
Elaboration conditions of the Co catalysts. T_f is the temperature of filaments.

Deposit	Nature	Catalyst deposition mode	Pressure (mbar)	Temperature (K)	Catalyst reduction	Temperature rate (K/min)	Power filaments (W) [T_f] (above 573K)	Plasma
					$C_2H_2:NH_3:H_2$ (%)			
Nanocones	Silicon oxynitride	Sputtering	UHV(573K) p 15	973	0:20:80	10	150 [2100 K]	Yes
Nanotubes	Carbon	Evaporation	UHV(973K) p 15	973	0:0:100	10	150 [2100 K]	No

Table 2
Elaboration conditions of the nanostructures by PE HF CCVD. For plasma conditions see SI-1.

Deposit	Nature	Pressure (mbar)	Temperature (K)	$C_2H_2:NH_3:H_2$ (%)	D_{flow} (sccm)	t (min)	Power filaments (W) [T_f]
Nanocones	Silicon oxynitride	15	973	20:16:64	100	15	150 [2100 K]
Nanotubes	Carbon	15	973	20:0:80	100	6	150 [2100 K]

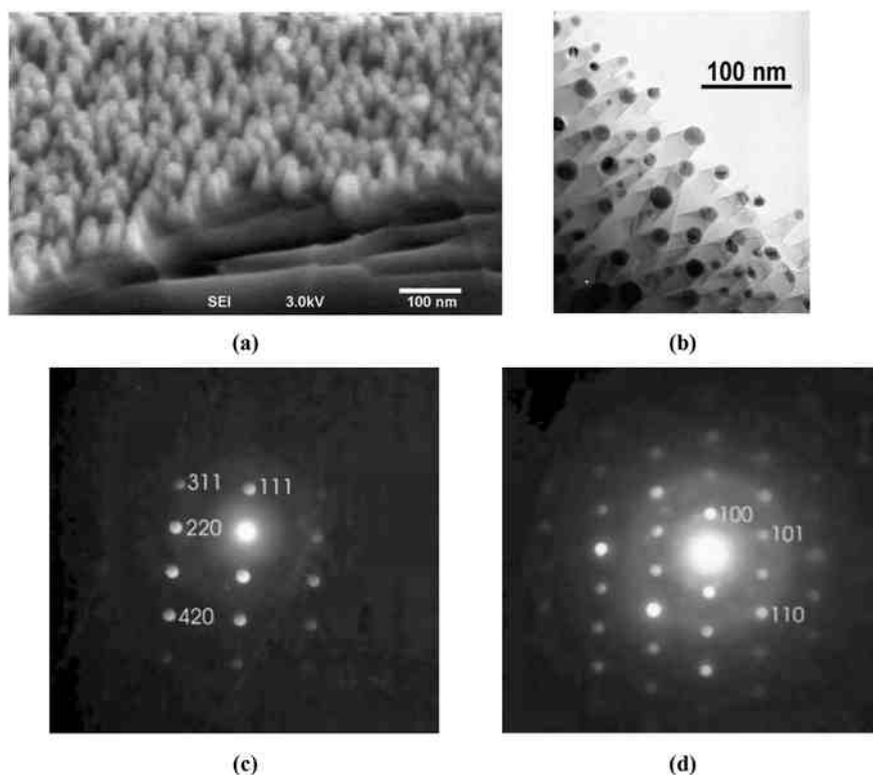


Fig. 1. SEM (a) and TEM (b) images of sample consisting of Co NPs on the top of Si oxynitride nanocones. Both images suggest aligned nanocones with a dark NP on top. Electron diffraction showing (c) cubic (fcc) and (d) hexagonal (hcp) lattices of Co NPs.

Supplementary Information 1 (SI 1) and 2 (SI 2), respectively. The size distribution is not completely monomodal with a main maximum at around 25 nm with secondary maxima around 40 nm and 13 nm, respectively. The density is around $4 \cdot 10^{10} \text{ cm}^{-2}$. The nanoparticles are isotropic. Morphology and geometrical parameters of Co NPs are summarized in Table 3.

The crystalline structure of the NPs has been studied by TEM electron diffraction. The Co was observed with both cubic fcc (Fig. 1c) and hexagonal hcp (Fig. 1d) lattices. The fcc and hcp patterns assignment are reported in Supplementary Information 3 (SI 3) together with the corresponding ASTM files. NPs were mainly fcc structure however.

Table 3
Morphology and geometry of the nanosized Co.

Deposit	Deposition process	Mean size (nm)	Density ($10^{10}/\text{cm}^2$)	NP's shape
Nanocones	DC magnetron (2.5 nm)	25 (40, 13)	4	Isotropic
Nanotubes	Evaporation (4.5 nm)	13	1.3	Anisotropic

The chemical nature of the nanocones was generally investigated by in situ X-Ray photoelectron spectroscopy (XPS) (non-monochromatized Mg K α source with VSW100 detector working at 0.65 eV resolution) or ex situ analyzed (monochromatic Al K α source with a VSW150 detector working at 0.55 eV resolution). Without exposure to air, the sample is transferred in a surface analysis chamber directly connected to the deposition/treatment chamber. In Fig. 2 we show the survey XPS spectra of a sample cleaned by the H $_2$:NH $_3$ plasma before (bottom spectrum, red line) and after the C $_2$ H $_2$:NH $_3$:H $_2$ exposition (upper curve, blue line) (i.e. before and after synthesis of nanocones with Co NPs on top). Both spectra were recorded with an unmonochromatized MgK α source. The presence of silicon (Si2p and Si2s), oxygen (O1s and O2s), nitrogen (N1s) and cobalt (Co2p and Co3p) on the surface is clearly evidenced. The most important effect is the presence of nitrogen, which indicates, together with the presence of oxygen, a pronounced plasma-enhanced CVD formation of silicon oxynitride (SiN $_x$ O $_y$). Their intensities do not much change with the two successive treatments. Importantly, only a negligible contribution of carbon is present (around 5% with graphitic nature at 284.6 eV), despite the acetylene injection into the CVD reactor. Also, the nitrogen concentration becomes even stronger after the C $_2$ H $_2$:NH $_3$:H $_2$ exposition. The individual spectra of the Co2p, Si2p, N1s and O1s core levels are reported in Supplementary Information 4 (SI 4) in two situations: either after reduction in UHV for Co2p line and after synthesis of nanocones for all of them. They are completed with Co2p core level spectrum after air removal followed with a UHV thermal annealing at 873K. All these spectra show the presence of a weak CoO contribution at 782.3 eV even after synthesis, besides the main Co 0 contribution at 777.8 eV.

The surface concentration extracted from the XPS Si2s, C1s, N1s, O1s, Co2p3/2 corelines are summarized in Table 4. It is seen that the main Si2p peak at 101.65 eV, the main N1s peak at 397.3 eV and the main O1s peak at around 532 eV can be associated with the silicon oxynitride (SiN $_{0.3}$ O $_{0.3}$) in good agreement with the XPS assignment reported on the surface oxidation of silicon nitride in the literature, with Si2p, N1s and O1s main peaks at 101.6, 397.6 and 532.7 eV, respectively [29]. The cobalt is mainly in a metallic form with a characteristic Co 0 2p3/2 contribution at 777.8 eV. The presence of some Co oxide (CoO) after the nanocones formation at 782.3 eV would mean that metallic Co on top of SiO $_x$ N $_y$ nanocone is covered

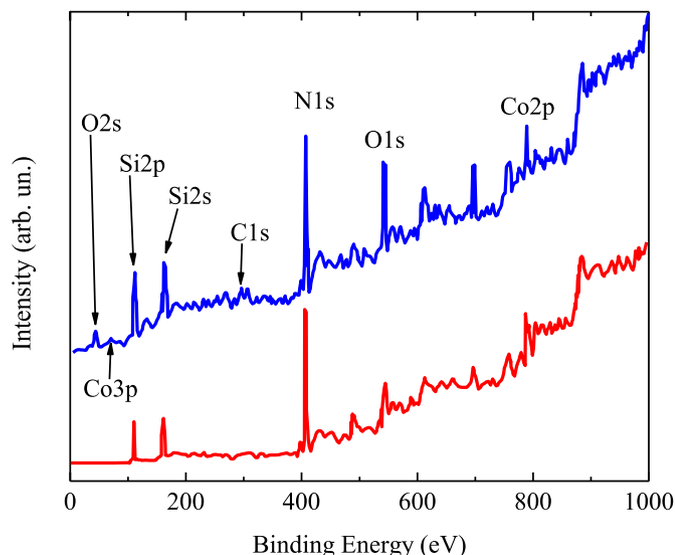


Fig. 2. XPS survey spectra before (bottom, red) and after (up, blue) synthesis of nanocones. In situ XPS analysis. (A colour version of this figure can be viewed online.)

with some oxide and thus represents some core/shell structure. Finally, the absence of carbon at the surface is confirmed by the very weak Raman signal in the carbon range (1100-1650 cm $^{-1}$) (not shown here).

2.2. Nanoparticles inside the CNT: synthesis and characterization

In the second approach, the catalyst was deposited with an evaporation cell in an UHV chamber directly connected to the synthesis chamber (4.5 nm of equivalent Co layer). The use of UHV conditions (base pressure 10 $^{-9}$ mbar) during the evaporation allows to finely control the metal dispersion as well as the metallic nature of the deposit. This is supported by the XPS spectra of wide scans before and after Co evaporation as well as by the Co2p core level reported in Fig. 3. The Co2p3/2 line exhibits a single line with a binding energy (777.8 eV) and an asymmetry of the line (with, according to a Doniach-Sunjić fitting analysis [30], a full width at half-maximum $G_{1/2}$ 0.6 eV and an asymmetry factor $a_{1/2}$ 0.37 instead of $G_{1/2}$ 0.61 and $a_{1/2}$ 0.375 [31]) quite coherent with metallic cobalt exclusively (Fig. 3d). The absence of impurity as well as the full metallic nature of evaporated Co render easier the reduction step with a single temperature rise at 973K at the rate of 10K/min.

The PE HF CCVD carbon nanotube growth was carried out with a mixture of C $_2$ H $_2$ and H $_2$ (without ammonia) at p $_{1/2}$ 15 mbar for 6 min. The conditions are summarized in Tables 1 and 2. These conditions (temperature, heating up and down, etc.) were chosen to optimize the formation of homogeneous distribution of NPs. In this case the substrate was not etched but a vertically aligned array of CNTs is formed [27,28]. Small time of synthesis was chosen to minimize the formation of defects in carbon nanotubes. The quality of the grown nanotubes has been checked by Raman spectroscopy (Renishaw apparatus with He-Ne laser source) (Fig. 4). The spectrum in the carbon domain exhibits weak D lines split at 1306 cm $^{-1}$ and at 1334 cm $^{-1}$, respectively. By contrast the intense G line exhibits a Breit-Wigner-Fano (BWF) shape with the main contribution at 1591 cm $^{-1}$ with a small full width at half maximum (FWHM) of 13 cm $^{-1}$ and weaker downward vibration modes at 1555 cm $^{-1}$, together with the D 0 mode at 1606 cm $^{-1}$. This spectrum strongly suggests that the carbon nanotubes are mainly conductive with a high quality. A very small contribution due to amorphous carbon at 1525 cm $^{-1}$ can be observed.

Further characterization of the samples has been done by atomic force microscopy (AFM: NANOSCAN Dimension 3100 used in the tapping mode with Si cantilever and lateral resolution 5e10 nm), SEM and TEM. These investigations strongly suggest the massive growth of vertically oriented nanotube (Fig. 5a). This is expected when using a plasma activated process [27]. Each CNT contains only one Co nanoparticle inserted and located at the top of the nanotube. The AFM image of the Co surface before CNT growth is presented in Fig. 6a and its 3D reconstruction is depicted in Fig. 6b, while the size distribution of the Co NPs is shown in Fig. 6c. The diameter distribution is well described by the log-normal distribution function as shown in Fig. 6c, with the average diameter around 13 nm. The density of NPs was evaluated to be 1.3 10 10 cm $^{-2}$ (Table 3). The TEM result confirms the presence of Co inclusions on the top of each CNT (Fig. 5b). The average length of CNT array is around 1 μ m. It should be emphasized that the major part of these Co nanoparticles is anisotropic, as can be seen in Fig. 5c and d, with the long axis always oriented parallel to the nanotube axis. Thus, in Fig. 5d the length L is around 70 nm and the revolution width W is 20e25 nm, which yields an aspect ratio L/W of around 3. Fig. 5c shows Co nanoparticles with an anisotropic nailhead form. This is the most characteristic feature of these samples as reported [32]. The diameter of these inclusions is restricted by the inner channel of carbon nanotube and their length can reach 250e300 nm, i.e. an

Table 4
Concentration analysis from the XPS data before and after the nanocones (nanotubes) formation.

Sample	Assignment	Si2p		N1s	O1s		Co2p _{3/2}		C1s
		Si	SiO _x N _y	SiO _x N _y	O _{surface}	SiO _x N _y (pCoO)	Co	CoO	
Before nanocone formation	Energy (eV) [Concentration, %]	100.8 [3]	101.6 [31]	397.4 [38]	535.0 [3]	533.1 [6]	778.1 [19]	e	e
After nanocone formation	Energy (eV) [Concentration, %]	e	101.65 [35]	397.3 [35]	533.6 [2]	531.65 [10]	778.1 [12]	782.6 [1]	284.6 [5]
After nanotube formation	Energy (eV) [Concentration, %]	e	102.3 [9]	e	e	533.6 [16]	777.9 [3]	e	284.6 [72]

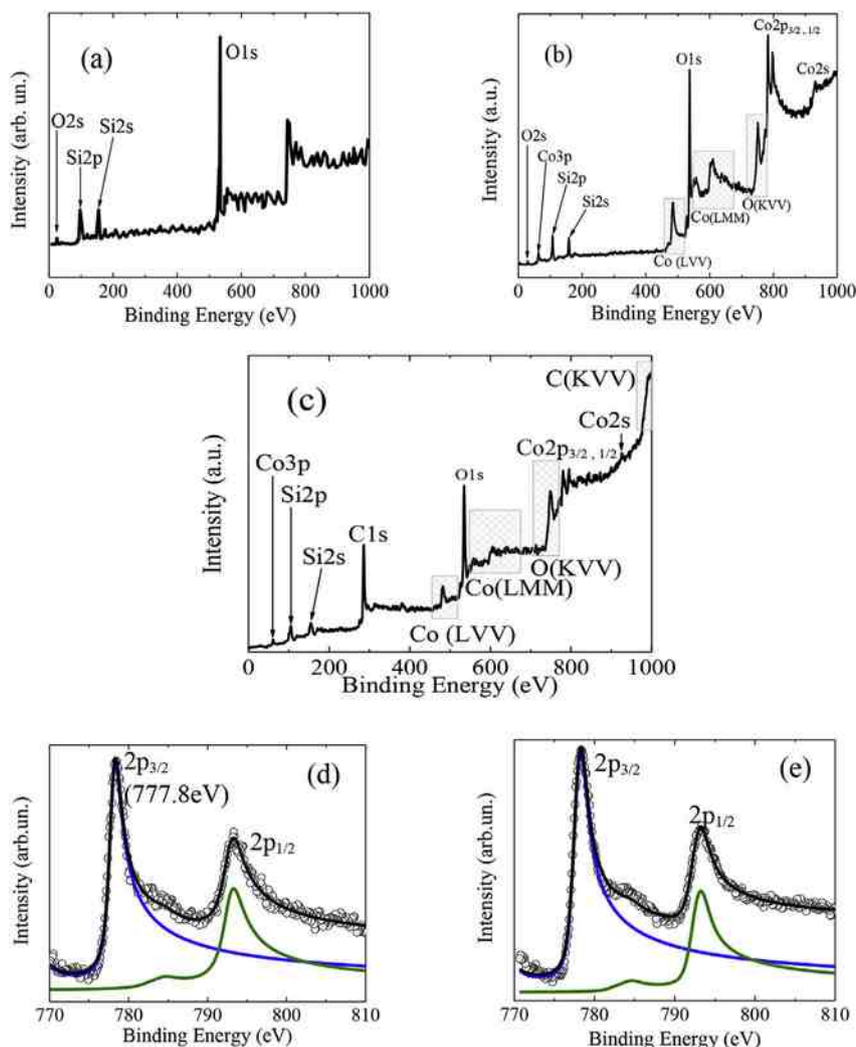


Fig. 3. (a) XPS wide scan on the SiO₂/Si(100) substrate before Co evaporation; (b) XPS wide scan and (d) Co2p core level after in situ Co evaporation; (c) XPS wide scan and (e) Co2p core level after carbon nanotube synthesis. We can note the absence of any carbon surface contamination before synthesis. In situ analysis with non-monochromatized Mg K α line. The Co 2p_{3/2} and Co 2p_{1/2} lines are fitted with two Doniach-Sunjić lines [30]. (A colour version of this figure can be viewed online.)

aspect ratio up to 20. Such drastic change of a morphology of catalytic Co NPs during the CCVD of carbon nanotubes confirms the complicated nature of the growth process. The most likely process occurs through a top growth mode described elsewhere [32]. We can exclude the possible melting of Co due to relatively low temperature of the CCVD (973K). However, the nailhead shape of the Co nanoparticle often observed suggests that some diffusion occurs inferring nanoparticle shape transformation. This can be explained by large incorporation of carbon atoms driving the formation into some viscous state at a temperature not far from the eutectic temperature of the cobalt nanoparticle, an intermediate state between the solid or liquid state. Anyway, such resulting Co morphology embedded on top of CNT underlines the importance of

diffusion processes in/on the NP bulk/surface during the CNT growth. TEM analysis reveal the regular structure of CNTs and their low defectiveness in agreement with the Raman spectroscopy. The nanotubes are multiwall but with a clear and regular hollow inside. More details about the processes of PE HF CCVD for vertically aligned CNT growth can be found elsewhere [27,28,32,33].

3. Magnetic properties

3.1. Magnetic force microscopy and FC-ZFC measurements

In Fig. 7 we show the results of the AFM and magnetic force microscopy (MFM) recorded on NANOSCAN Dimension 3100 of

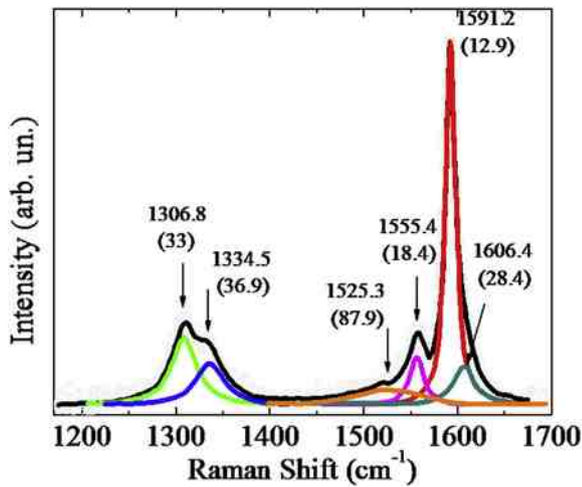


Fig. 4. Raman spectrum of arrays of vertically aligned CNT acquired with the laser wavelength of 632.8 nm. Numbers indicate the Raman shift value for a peculiar mode. The numbers in brackets indicate the full width at half maximum in the same mode. Points are for the experimental data. Colored lines correspond to the different vibration modes specified in the text. The black line is the envelope obtained from all these considered modes. (A colour version of this figure can be viewed online.)

arrays of Co NPs at the top of silicon oxynitride nanocones. Sample was previously without magnetic history and both AFM and MFM images were acquired on the same part of the sample. To separate the short-range Van der Waals forces from the long-range magnetic forces, the same area has been scanned twice at distance z (AFM image) and z_{pd} , successively. The distance between the sample and cantilever was taken as $d \approx 500$ nm. Then the contribution of the magnetic force is determined from the phase variation which is proportional to the second derivative of the magnetic field. Despite the very rough character of the surface it is possible to detect by AFM the nanocones, with a size lower than 100 nm. Moreover, by MFM it is possible to detect magnetic domains on the surface of the sample with average sizes around 500 nm encompassing a set of

many nanocones, which confirms the strong DDI between Co NPs. Other images are given in SI-5. It should be emphasized that these images were obtained on the sample without magnetic history, but we failed to record contrasted MFM images on the sample with a remanent magnetism after exposition to a magnetic field parallel or normal to the surface. This is likely due, as will be seen further, to the low coercive field ($H_c \approx 140$ Oe) that can be reversed by the magnetic field of the cantilever in the former case, and to the very low remanent magnetization $M_r \approx 0.07 M_s$ where M_s is the magnetization at saturation in the latter case.

In Fig. 8a we show the FC-ZFC magnetization curves for the same sample measured using a MPMS SQUID-VSM magnetometer (Quantum Design) combining the high speed of a VSM (vibrating sample magnetometer) with the sensitivity of the SQUID (Superconducting Quantum Interference Detector) magnetometer (MPMS). The magnetic field $H \approx 25$ Oe was oriented parallel to the surface substrate. The characteristic feature of the FC-ZFC curve is that up till 390 K the system never reaches the equilibrium state at which the field cooled magnetization, M_{FC} becomes equal to zero-field cooled magnetization, M_{ZFC} . Blocking temperature T_B for the investigated Co NPs is close to 400 K. It means that below this value nanoparticles are ferromagnetic with well-defined coercivity (not shown here). Importantly, the $M_{FC}(T)$ curve is almost temperature-independent for $T < 400$ K. This is again an indication of strong DDI between magnetic NPs [37,38]. Thus, two independent measurement techniques confirm the presence of a strong DDI between the nanoparticles localized at the top of SiO_xN_y nanocones.

Now we move to the AFM-MFM and FC-ZFC magnetization results for the Co NPs localized inside CNT. In Fig. 9 AFM-MFM images are displayed for this sample. It is clearly seen that NPs are magnetically decoupled. It follows from the presented result that there is a unique correlation between the geometrical sizes of particles and the size of magnetic domains, - each particle corresponds to a single domain.

In Fig. 8b we present the M_{FC} and M_{ZFC} plots versus temperature measured at $H \approx 25$ Oe for arrays of Co NPs embedded inside CNTs. Magnetic field was applied parallel to the substrate surface. The blocking temperature is around 400 K, like for Co NPs on silicon

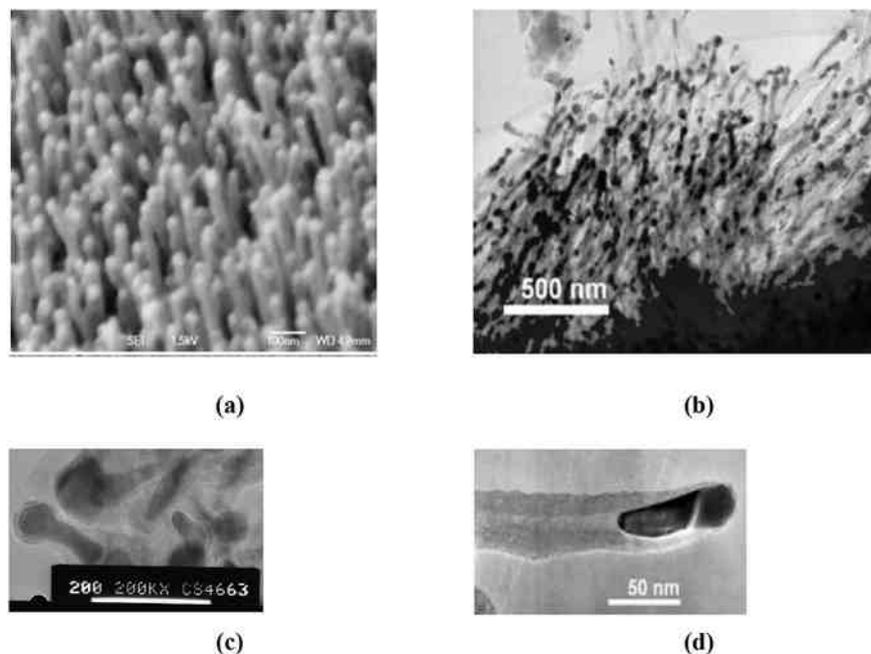


Fig. 5. (a) SEM and (b) TEM images of CNT grown on SiO_2/Si substrate with Co NPs; (c) and (d) display the structural shape of the Co nanoparticles.

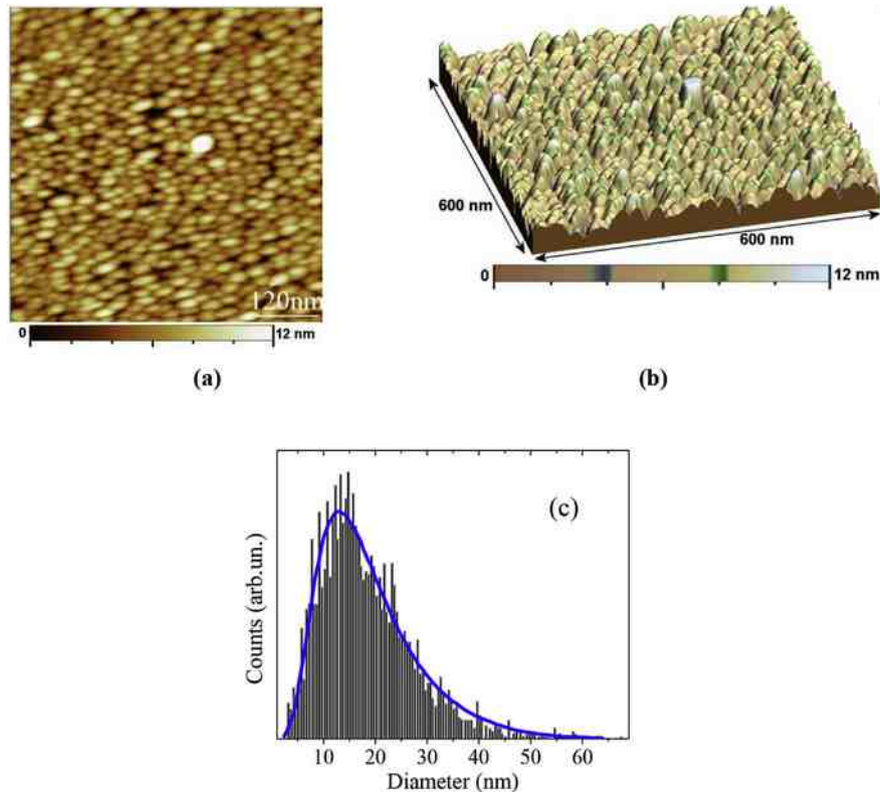


Fig. 6. (a) AFM image of Co NPs on SiO₂/Si substrate; (b) Three-dimensional reconstruction of the same surface; (c) Distribution of the Co NP's size histogram (5000 nanoparticles) fitted with a log-normal law (blue curve) peaking at around 13 nm. (A colour version of this figure can be viewed online.)

oxynitride nanocones. Unlike the previous sample however, the $M_{FD}(T)$ behavior is quite different. Now it almost linearly increases when temperature goes to zero. This is characteristic feature of magnetically decoupled NPs [4,11,34,35].

In conclusion, from the data presented in this section, it follows that the competition between the DDI and intrinsic properties of ensemble of Co NPs of approximately of the same diameter and the same density distribution varies drastically according to the preparation process. When NPs are localized on the top of silicon oxynitride nanocones, the DDI dominates. When NPs are embedded inside CNTs, the intrinsic interaction becomes dominant. This is a major result of this comparative work. In the following, we will try to elucidate why the DDI between Co inclusions becomes less dominant and explore the role of CNT in this.

3.2. Analysis of the M(H) loops

Important information can be obtained from the analysis of the hysteresis loops $M(H)$. The measured $M(H)$ loops for samples Co-SiO_xN_y and Co-CNT at T ¼ 2K are presented in Fig. 10a and Fig. 10b, respectively. Data for the magnetization are normalized to the saturation value M_s . Similar results are displayed in SI-6 for T ¼ 350 K. Let us recall that the parallel and perpendicular magnetic field orientations were determined relative to the plane of the substrate. For sample Co-SiO_xN_y in parallel field the magnetization saturates very fast, the saturation field is close to 3 kOe. On the contrary, in the perpendicular field, the saturation field is much greater and exceeds 20 kOe. The coercivity H_c and the squareness M_r/M_s in the parallel field is equal to 820 Oe and 0.56, correspondingly. In the perpendicular field these quantities quote to 414 Oe and 0.092, respectively. Rapid saturation in the parallel field along with greater values of the squareness and the coercivity can

be reasonably attributed to an easy axis of magnetization which is oriented parallel to the surface of the substrate. This can be naturally explained by the strong DDI between NPs.

Reversely, for Co-CNT sample $M(H)$ loops approach each other for a parallel and perpendicular field directions (Fig. 10b). In the parallel field, i.e. perpendicular to the nanotube axis, a slightly faster saturation is observed, akin to the magnetization along the easy axis. In the direction of the magnetic field perpendicular to the substrate, i.e. parallel to the nanocylinder axis, the approaching saturation is slightly slower, but the magnetization curve is much closer to the case of the parallel field, compared to, for example, the magnetization of Co nanowires along their easy axes [30]. In Ref. 36 curves of approaching magnetic saturation along easy and hard axes, as well as coercivity and anisotropy constants differ significantly.

Thus, an essential difference in the shape of the magnetization curves is observed when nanosized Co is formed on silicon oxynitride and inside CNT, respectively. In the former case the easy axis of magnetization can be easily identified as oriented in-plane, while in the latter case axes of the Co nanocylinders are neither easy nor hard axes in a first approach. The result obtained for Co-CNT sample is not typical for systems of Co nanocylinders (nanowires). Usually in such systems, with aspect ratios equal to 3 and greater, the shape anisotropy dominates, and easy axis is oriented along the cylinder axis and the $M(H)$ shapes for easy and hard axes directions are very different [36e39]. In our samples the experimental data reveal that this is not the case. Although $M(H)$ curves slightly differ, they are much closer to each other than usually reported in literature [36e39]. Therefore, most likely, here is a complex case where magnetocrystalline and, possibly, magnetoelastic anisotropy of nanoparticles have a dispersion relative to some direction at a certain angle to the axis of the nanocylinders.

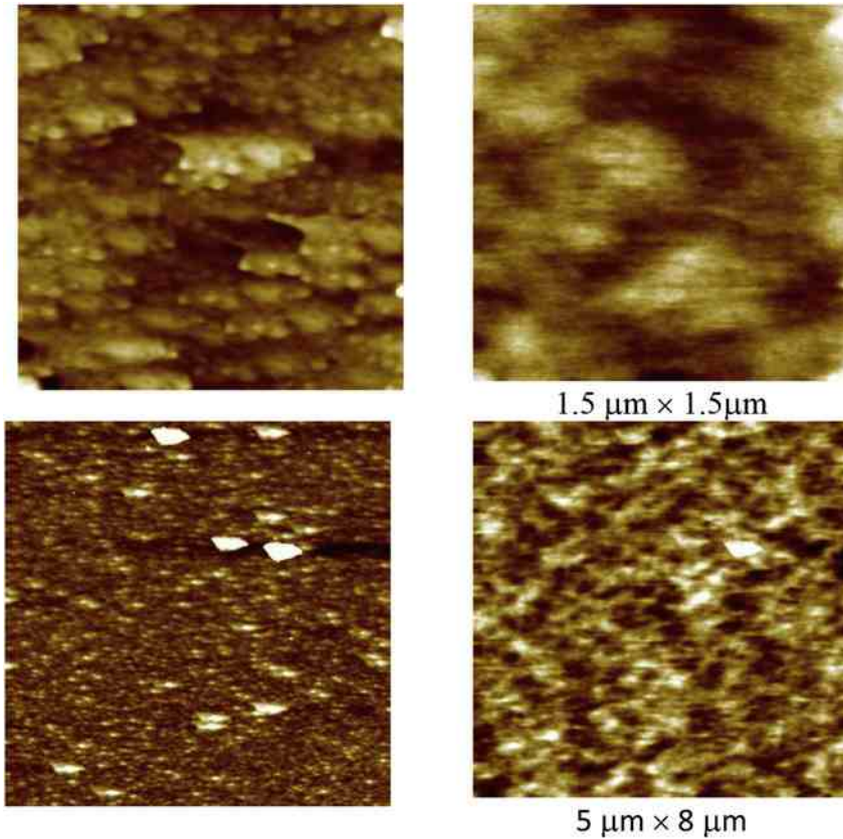


Fig. 7. AFM (left) and MFM (right) images of arrays of Co NPs on the top of silicon oxynitride nanocones. Sample without magnetic history. (A colour version of this figure can be viewed online.)

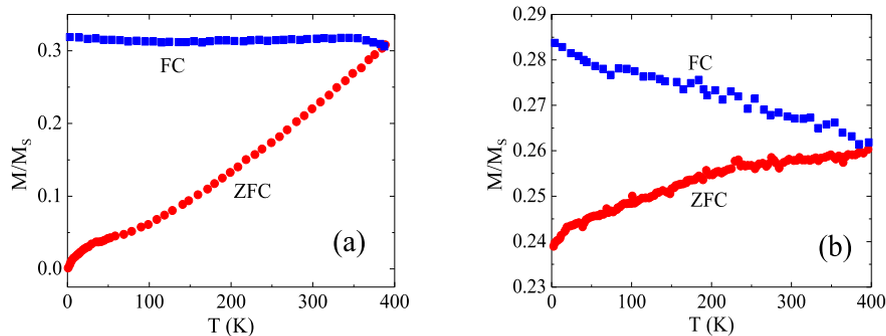


Fig. 8. FC-ZFC magnetizations for (a) Co NPs on the top of silicon oxynitride nanocones; (b) Array of Co NPs embedded inside CNTs. Magnetization was measured at $H \parallel 25$ Oe. (A colour version of this figure can be viewed online.)

From these results it follows that, the hexagonal c axis of hcp Co does not correlate with the nanocylinder axis as is usually observed for ensemble of Co nanowires [39]. Therefore, the magnetocrystalline anisotropy is neither oriented along the nanocylinder axis nor perpendicular to it. Both the magnetocrystalline anisotropy of the material and the shape anisotropy of nanoparticles embedded inside CNT have a competing effect on the magnetic properties of Co-CNT nanocomposite in this case. The tilt of c axis of hcp Co could be due to the peculiar mechanism of nanosize Co fabrication. Indeed, the formation of an array of cobalt nanoparticles occurs by depositing first a nanosized film on amorphous silicon oxide substrate. At the stage of nucleation, the Co film deposited on a non-oriented substrate does not wet the surface and Co crystallites aggregate with random crystalline orientations

[40,41]. The insertion of the NPs inside carbon nanotubes could create stresses and deformation that, in turn, could be the reason of the occurrence of a magnetoelastic anisotropy.

4. Discussion

4.1. Evaluation of the micromagnetic parameters

In this Section we will evaluate effective micromagnetic parameters like anisotropy (H_{ra}) and exchange (H_{ex}) fields, and the effective magnetic anisotropy constant (K). All these quantities are evaluated within the random anisotropy model (RAM) [42,44]. The RAM has been successfully applied in the past to explain the properties of amorphous and nanocrystalline ferromagnets [44,45],

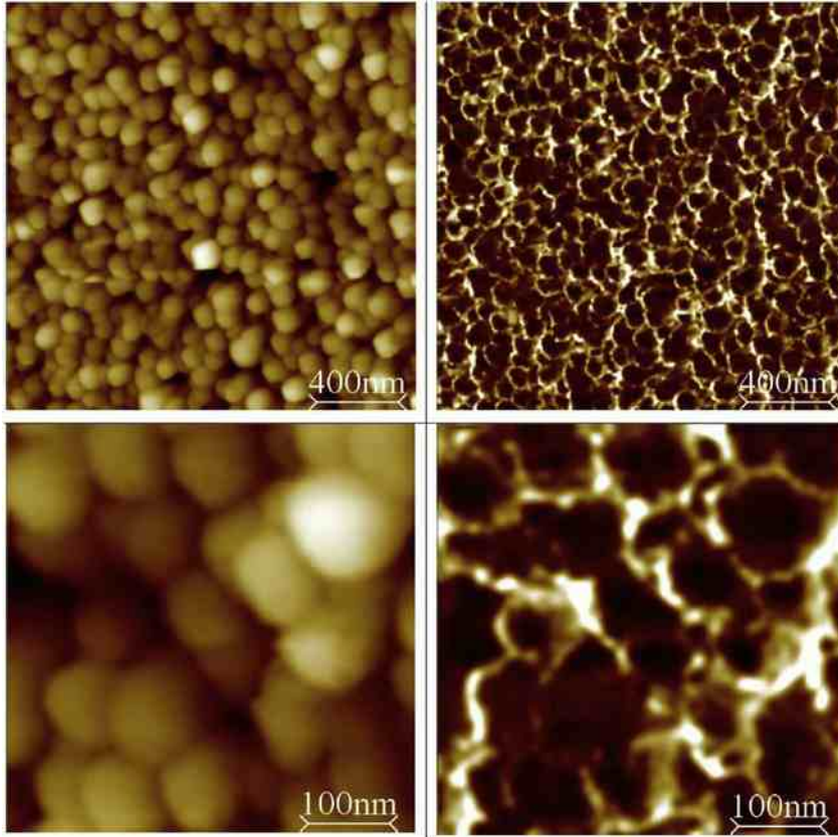


Fig. 9. AFM (left) and MFM (right) images of arrays of Co NPs embedded inside CNTs at different magnifications. (A colour version of this figure can be viewed online.)

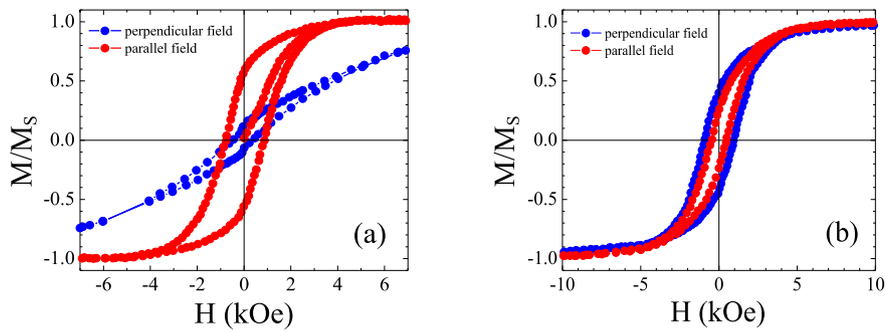


Fig. 10. $M(H)$ curves for the parallel (red color) and perpendicular (blue color) magnetic field orientations. $T = 2$ K (a) sample Co-SiO_xN_y; (b) sample Co-CNT. (A colour version of this figure can be viewed online.)

and more recently to explain the properties of ferromagnetic NPs in CNT [21,24,46,47].

The basic equations of the RAM are reduced to the following simple expressions for the micromagnetic parameters [21,24,43].

$$H_{ra} \approx \frac{2K}{M_S} \quad (1)$$

$$H_{ex} \approx \frac{2A}{M_S R_a^2} \quad (2)$$

$$K \approx \frac{h}{4} H_c^3 M_S - 114 \delta k_B T_B^2 \quad (3)$$

where A is the exchange constant, k_B is Boltzmann constant, and R_a is the length over which the magnetic anisotropy axes are correlated. Usually in a nanocrystalline material R_a is assumed to be

equal to the radius of the nanoparticle [21,44].

From Eqs. (1)-(3), it follows that in order to evaluate the micromagnetic parameters it is necessary to measure a number of quantities such as M_S , T_B , H_c , R_a and to know the exchange constant A . The T_B , H_c , and R_a were obtained by us directly from the experiments. The M_S could also be obtained from the experiment if the exact weight of Co is known and under the assumption that all the cobalt NPs are ferromagnetic. The weight of Co was estimated based on the average density of the nanoparticle arrangement and their average size. The obtained value was consistent with the amount of evaporated Co according to Table 1. Nevertheless, some uncontrollable distribution of quantities used in such calculations can lead to significant inaccuracies in the M_S value. For this reason, we also analyzed the literature data for M_S in nanosized Co samples. The A values were also taken from literature data for the ensemble

of Co NPs.

It is known that the saturation magnetization for nanosized Co is characterized by significant dependencies on the temperature and size of the NPs [48,49]. In the low-temperature region its value for clusters containing more than 550e600 atoms approximates the value of a bulk material [50], for which $M_S \approx 1440 \text{ emu/cm}^3$ [51]. Ross et al. assumed that $M_S \approx 1100 \text{ emu/cm}^3$ for massive of Co NPs [52]. To calculate the magnetization distribution in Co NPs, it is usually assumed that the saturation magnetization of Co NPs is $M_S \approx 1400 \text{ emu/cm}^3$, the exchange constant is given by $A \approx 1.3 \cdot 10^6 \text{ erg/cm}$ and the uniaxial anisotropy constant is $K \approx 4.3 \cdot 10^6 \text{ erg/cm}^3$ [53,54]. As for the exchange constant A, it depends less on temperature, but for nanoscale layers or nanoparticle arrays its value may depend on the method by which it is measured. Measurements of A indicate a large discrepancy in the values: $1.8 \pm 0.3 \cdot 10^{11} \text{ J/m}$ from Brillouin light scattering (BLS) [55], $2.7 \pm 0.1 \cdot 10^{11} \text{ J/m}$ from neutron scattering [56], and $2.1 \cdot 10^{11} \text{ J/m}$ from ferromagnetic resonance studies [57]. It was established in Ref. [58] from BLS data that A and M_S of the Co layer of 80 nm are equal to $2.1 \cdot 10^{11} \text{ J/m}$ and 1332 emu/cm^3 , respectively, and for 10 nm thickness A was found to be $1.54 \pm 0.12 \cdot 10^{11} \text{ J/m}$ and M_S was $1247 \pm 44 \text{ emu/cm}^3$.

Based on the experimental data and analysis of the literature, we evaluated H_{ra} and H_{ex} values together with K. All measured and evaluated data are summarized in Table 5 for Co-CNT and Co-SiO_xN_y systems.

It follows from the analysis for both samples that the exchange field is less than the anisotropy field, which could indicate the negligible role of the interparticle exchange coupling. It is also confirmed by relatively high values of the anisotropy constant, $(2.65e3.77) \cdot 10^5 \text{ J/m}^3$ and random anisotropy field, $(3.76e5.35) \text{ kOe}$, which could be due to shape, magnetocrystalline and magnetoelastic anisotropies. In the above considerations the DDI was absent. Obviously, for densely packed NPs its contribution could be essential. These arguments require further analysis.

4.2. Comparative analysis of the contributions of SA and DDI

The total anisotropy energy is determined by the contributions of the magnetocrystalline (K_{MCA}) and shape anisotropy (K_{SA}), the magnetoelastic contribution (K_{MEA}), the DDI (K_{DD}) and possibly the surface anisotropy. We will neglect this last term further in this study. The effective anisotropy constant K found above, determines the total contribution of all components [60,61].

$$K \approx K_{MCA} + K_{SA} + K_{DD} + K_{MEA} \quad (4)$$

The shape anisotropy is determined by the interaction of the particle with its own demagnetizing field. For a homogeneous magnetization the corresponding SA constant is [54].

$$K_{SA} \approx \delta_1 = 2\mu_0 M_S^2 DN; \quad (5)$$

where $DN \approx N_{\parallel} - N_{\perp}$, N_{\parallel} is the demagnetizing factor of the NP along the symmetry axis z and N_{\perp} is the demagnetizing factor in the direction perpendicular to z axis.

The contribution of the DDI between NPs is expressed as [62,63].

$$K_{DD} \approx \delta_1 = 8\mu_0 \mu_0 M_S^2 V^{-1} z^3 s_d; \quad (6)$$

where z is an average interparticle distance, s_d is a lattice sum that depends on the type of lattice in the NPs arrangement, their shape, on the dimensionality of the sample and on the mutual orientation of the NPs magnetizations, V is the NP volume.

The contribution of the SA for the sample Co-SiO_xN_y can be neglected due to the almost equal transverse and longitudinal demagnetizing factors ($DN \ll P$), P being the volume fraction occupied by the ferromagnetic material. Due to the spherical shape of NPs in this case it seems reasonable to also neglect the possible contribution of the magnetoelastic anisotropy. Therefore, the anisotropy constant K in this case is a function of only K_{DD} and K_{MCA} . To estimate the contribution of the DDI, we follow Eq. (6), in which for two-dimensional arrays of NPs the value of the lattice sum $s_d \approx 6e8$ was applied [64,65].

For the arrays of Co-CNT, we cannot neglect the possible contribution of SA, MCA and MEA. For these samples the total anisotropy constant will be determined by the expression (4). The contribution of the SA and DDI of the system of magnetic nanocylinders is expressed as [66].

$$K_{SA} + K_{DD} \approx \delta_1 = 4\mu_0 M_S^2 \delta_1 \cdot 3P; \quad (7)$$

here $P \approx (\pi r^2 S)$ is a volume fraction occupied by ferromagnetic nanocylinders, r is radius of the nanocylinder, S is surface density of the nanocylinders. In Table 6 we present the results of the evaluations of the SA, MCA and DDI for both samples for the parallel field. We do not show K_{MEA} in this Table. The discussion of this issue will be done in the next Section.

Results presented in Table 6 reveal that in the case of Co-SiN_xO_y array, in which the NP shape is close to spherical, K_{MCA} is slightly greater than K_{DD} . Nevertheless, the impact of the DDI in the presence of only MCA is determined by the dipolar coupling constant $a \approx K_{DD} = K_{MCA}$ [67]. There is a crucial parameter a_{cr} that determines the transition from single-particle to collective behavior. The reported values for fcc or hcp Co NPs are $a_{cr} \approx (0.2e0.4)(d/z)^3$ [68]. If $a > a_{cr}$, then DDI dominates, otherwise the MCA prevails. For our geometry we get $a_{cr} \approx 0.14 < a \approx 0.61$, which indicates that the contribution of the DDI is sufficient for the formation of regions covering many NPs, that is indeed observed experimentally.

For the Co-CNT array we have to consider both the SA and MEA.

Table 5
Micromagnetic parameters of Co-CNT and Co-SiO_xN_y systems as obtained within RAM.

Parameter	Unit	Parallel field		Perpendicular field		Reference note
		Co-CNT	Co-SiO _x N _y	Co-CNT	Co-SiO _x N _y	
H_c	Oe	473	820	1050		Experiment
M_S	A/m	$1.4 \cdot 10^6$	$1.4 \cdot 10^6$	$1.4 \cdot 10^6$	$1.4 \cdot 10^6$	Experiment and Ref. [53,54]
A	J/m	$1.54 \cdot 10^{11}$	$1.54 \cdot 10^{11}$	$1.54 \cdot 10^{11}$	$1.54 \cdot 10^{11}$	Ref. [59]
T_B	K	400	400	~500	314	Experiment
M_r/M_S		0.25	0.25	0.45	0.45	Experiment
K	J/m ³	$2.65 \cdot 10^5$	$3.77 \cdot 10^5$	$3.15 \cdot 10^5$	$3.15 \cdot 10^5$	Calculations
H_{ra}	kOe	3.76	5.35	4.48	4.48	Calculations
H_{ex}	kOe	1.4	2.19	1.4	2.19	Calculations

Table 6
Evaluations of the SA, MCA and DDI for the parallel orientation of the magnetic field. d is an average diameter of the NP and L is an average height of the nanocylinder.

Parameter	Unit	Co-SiO _x N _y	Co-CNT
K_{SA}	J/m ³	e	6.16 10 ⁻⁵
K_{DD}	J/m ³	1.4 10 ⁻⁵	1.09 10 ⁻⁵
K_{MCA}	J/m ³	2.37 10 ⁻⁵	2.37 10 ⁻⁵
$K_{SA} \beta K_{DD}$	J/m ³	e	5.07 10 ⁻⁵
$a \frac{1}{4} K_{DD}/K_{MCA}$	a.u.	0.61	e
Z	nm	32.6	51.5
S	cm ²	3 10 ⁻¹⁰	1.2 10 ⁻¹⁰
P	a.u.	0.15	0.06
d	nm	25 ± 8	15 ± 5
L	nm	e	100e150

Their contribution along with the MCA should exceed the contribution of the dipole interaction between NPs.

4.3. Contribution of the magnetoelastic anisotropy (MEA)

Let us estimate the possible contribution of elastic stresses to the energy of magnetic anisotropy in Co-CNT samples. The total anisotropy energy could be determined by Eq. (4) [60,61]. All constants in Eq. (4) except the magnetoelastic anisotropy, have been already determined and given in Table 6. We reasonably assume that the K_{MCA} evaluated for the Co-SiO_xN_y samples will be the same as for Co-CNT. On the base of these data we obtained $K_{MEA} \approx 5.35 \cdot 10^{-5} \text{ J/m}^3$.

It should be noted that the type of crystalline lattice of Co affects significantly the interpretation of data on the magnetoelastic anisotropy [69]. For fcc Co the constant of the magnetoelastic anisotropy can be evaluated as

$$K_{MEA}^{fcc} \approx \frac{3}{2} \lambda \sigma \quad (8)$$

where λ is the magnetostriction constant and σ is the elastic stress [70]. Using for fcc Co $\lambda \approx 50 \cdot 10^{-6}$ [71] and applying the obtained K_{MEA} value we get σ order of 7.13 GPa.

For hcp Co when the hexagonal axis [0001] is parallel to the CNT axis and neglecting the torsional stresses, the constant of the magnetoelastic anisotropy can be expressed as [72].

$$K_{MEA}^{hcp} \approx \frac{c_{11} \beta c_{12}}{c_{33}} \frac{2c_2^2}{c_3} \left(\lambda_A \beta \lambda_B \beta \frac{1}{4} \sigma \lambda_A \beta \lambda_B \beta \right) \quad (9)$$

where c_{ij} are elastic stiffness constants, the indices $i, j \in \{1, 2, 3\}$ number the axes of the hexagonal crystal (here indices 3 corresponds to the hexagonal axis [0001]), λ_A and λ_B are magnetostriction constants and ε_1 is strain. Substituting in Eq. (9) $\lambda_A \approx 50 \cdot 10^{-6}$, $\lambda_B \approx 107 \cdot 10^{-6}$ [73], $c_{11} \approx 307.1 \text{ GPa}$; $c_{12} \approx 165.0 \text{ GPa}$; $c_{33} \approx 102.7 \text{ GPa}$; $c_2 \approx 358.1 \text{ GPa}$ [74] and using the previously obtained $K_{MEA}^{hcp} \approx 5.35 \cdot 10^{-5} \text{ J/m}^3$, we get $\varepsilon_1 \approx 0.00825$ which leads to the tensile stresses $\sigma \approx 3.408 \text{ GPa}$.

For the hexagonal axis 3 oriented perpendicular to the CNT axis the exact expression for the K_{MEA}^{hcp} depends on the orientation of crystal magnetization m [69],

$$K_{MEA}^{hcp} \approx \frac{1}{2} g B_1 \varepsilon_1 \beta B_2 \varepsilon_3 \beta B_3 \alpha_1 \beta \varepsilon_2 \beta \quad (10)$$

where $g \approx a_1^2 \beta \frac{\varepsilon_2}{\varepsilon_1} a_2^2 - \alpha_1^2 \beta a_2^2 \beta$ and a_i are the direction cosines of the magnetization with respect to the hexagonal axes, B_i are magnetoelastic coupling coefficients. In these calculations we used

$B_1 \approx 8.1 \cdot 10^{-6} \text{ J/m}^3$, $B_2 \approx 29 \cdot 10^{-6} \text{ J/m}^3$, $B_3 \approx 28.2 \cdot 10^{-6} \text{ J/m}^3$ [69]. The strain ε is found from the system of equations.

$$\begin{pmatrix} 0 & 1 & 0 & 1 \\ c_{11} & c_{12} & c_{13} & \varepsilon_1 \\ c_{12} & c_{11} & c_{13} & \varepsilon_2 \\ c_{13} & c_{13} & c_{33} & \varepsilon_3 \end{pmatrix} \begin{pmatrix} S \\ A \\ A \\ S \end{pmatrix} = \begin{pmatrix} 0 \\ 0 \\ 0 \\ 0 \end{pmatrix} \quad (11)$$

If axis 1 is oriented perpendicular and axis 2 parallel to the CNT axis, then we get from Eq. (11) $\varepsilon_1 \approx 0.00403S$, $\varepsilon_2 \approx 0.00300S$, $\varepsilon_3 \approx 0.00250S$, where S is in GPa. And for limiting cases $a_1 \approx 0$ and $a_2 \approx 0$ the value of g varies within the range $3/4 \dots 1$. This leads to boundary values of stresses from $S \approx 5.42 \text{ GPa}$ to $S \approx 21.93 \text{ GPa}$. The real value of stresses belongs to this interval. A similar result is obtained when the axis 3 and the axis 2 are perpendicular and the axis 1 is parallel to the axis of the CNT. However, the exact values of the direction cosines a_i are not known in our case. Therefore, for an approximate estimation of K_{MEA}^{hcp} we assume that the first term in Eq. (10) can be neglected and Eq. (10) can be rewritten as $K_{MEA}^{hcp} \approx \frac{1}{2} B_2 \varepsilon_3 \beta B_3 \varepsilon_1 \beta \varepsilon_2 \beta$. In this case the value of stress is $S \approx 9.515 \text{ GPa}$. This is valid for both cases, when the axis 3 is perpendicular to the axis of the CNT, and the axis 1 is either perpendicular or parallel to the axis of the CNT.

The elastic stiffness constants c_{ij} , generally speaking, depend on the stress S . This dependence is close to linear, $c_{ij}(S) \approx (1 + kS)c_{ij0}$, where c_{ij0} are the table values for stresses close to zero and $k \approx 0.017 \text{ GPa}^{-1}$ is an empirical coefficient [75,76]. Thus, considering the dependence of the magnetoelastic constants on the compression value, we have $S \approx 11.351 \text{ GPa}$. Such internal elastic stresses could cause the deformation of the CNT lattice only in the range of 0.5e2%, which is in good agreement with the literature [77].

The obtained K_{MEA} values reveal that the largest contribution of the magnetoelastic anisotropy occurs in the direction parallel to the substrate and when the hexagonal [0001] axes is oriented along the field direction. Therefore, we may conclude that the DDI for Co-CNT samples is overcome by the shape and magnetoelastic anisotropies. In the latter case the contribution of the hcp Co dominates. The c axes of the hcp phase with high probability is oriented perpendicular to the CNT axis. This assumption has been confirmed by the results of micromagnetic simulation, which was performed for hcp Co nanocylinders with diameter 20 nm and length 100 nm. We applied the Nmag package [78] based on the Landau-Lifshitz-Gilbert equation and finite elements method. The discretization length was chosen to be 1.5 nm that should be enough for the acceptable accuracy because the exchange length of cobalt is about 10 nm [79]. Axis Z is oriented along the nanocylinder axis (i.e., the CNT orientation), and XY plane is lying in the direction perpendicular to the CNT axis. Moreover, the X axis coincides with the [0001] axis of the hcp Co.

The simulation shows that in such conditions the relaxed magnetization configuration of the cylinder depends on the K_{MEA} (see Fig. 11). For the absence of the magnetoelastic stress, the magnetization of the Co nanocylinder is homogeneous and oriented along the Z axis, see Fig. 11a. The presence of the magnetoelastic stresses, $K_{MEA} \approx 5.35 \cdot 10^{-5} \text{ J/m}^3$, breaks nanocylinder on two domains, Fig. 11b. The orientation of the magnetization within the domain is oriented along the X axis. The performed simulations of the magnetic structure of Co nanocylinders should be considered only as evaluation results showing that with the magnetoelastic component nanocylinders may be partitioned into domains, which leads to a decrease in remanence. This correspond well to low remanence in Co-CNT samples. For hcp Co with c axis oriented along the axis of CNT, the reasonable values of K_{MEA}^{hcp} do not create the multidomain structure (this result is not shown here).

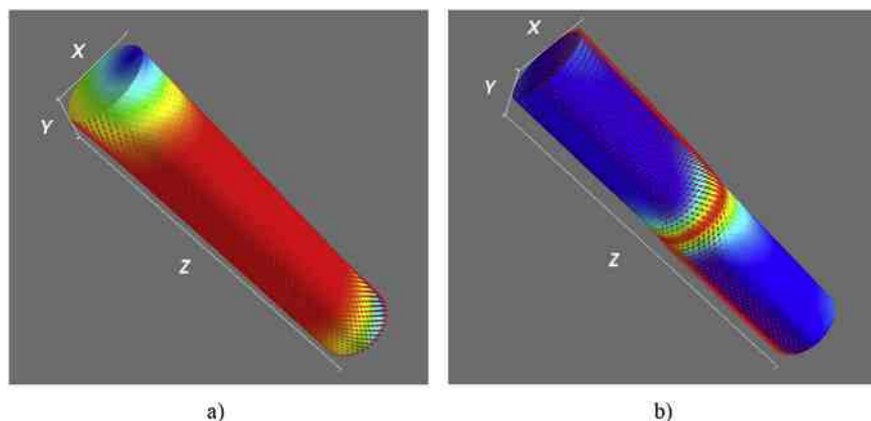


Fig. 11. Relaxed magnetization configuration of hcp Co nanocylinder of diameter 20 nm and length 100 nm for the absence of stress, $K_{\text{MEA}} = 0$ (a) and $K_{\text{MEA}} = 5.35 \cdot 10^{-5} \text{ J/m}^3$ (b). Red color corresponds to the magnetization direction along the nanocylinder axis, blue color corresponds to the magnetization direction oriented perpendicular to the nanocylinder axis. (A colour version of this figure can be viewed online.)

5. Conclusion

The analysis of the magnetic properties of nanosized Co on the top of SiO_xN_y nanocones and inside CNT has been performed. The average density of Co inclusions is similar for both type of samples and is order of 10^{10} cm^{-2} . Nevertheless, these two systems behave differently in a magnetic field. In the former case the Co NPs are strongly coupled via the DDI. As a result, the easy axis plane is oriented parallel to the substrate. For Co-CNT samples Co nanoparticles are magnetically isolated. The reason for a minor effect of the DDI is the magnetic anisotropy. It increases significantly because of the peculiar morphology of Co inclusions in CNT and stresses caused by the carbon nanotubes. We evaluated the values of shape, magnetocrystalline and magnetoelastic anisotropy constants. The magnetoelastic anisotropy has been evaluated for both observed crystalline structures of Co, fcc and hcp. It has been demonstrated that due to the presence of the hcp crystalline phase, stresses could reach significant values leading to noticeable magnetoelasticity. The influence of stresses onto magnetic structure of Co inclusions has been investigated by the micromagnetic simulations, which helped us to determine the probable direction of the hexagonal axis oriented perpendicular to the CNT axis. The developed approach in the fabrication of an ensemble of anisotropic nanoparticles can be very useful in the design of carbon-based magneto-electronic devices.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <https://doi.org/10.1016/j.carbon.2018.08.024>.

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