Very Low Potential Electrodeposition of Sm-Co Nanostuctures in Aqueous Medium Using Hard Templates

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Arrays of SmCo nanowires (NW) and nanotubes (NT) with a diameter of 200 nm and different lengths are synthesized by electrodeposition into the nanopores of an alumina membrane. The potential applied during the synthesis largely determines the nanostructure morphology, its crystallinity and composition. Potentials investigated are between −0.8 V and −3.0 V; in the potential range between −0.8 V and 1.0 V, long and perfectly ordered nanowires are obtained with a composition close to that of the equilibrium Sm2Co17 phase in the binary alloy. For higher potentials, above −1 V, samples are nanotubes, 195 nm in external diameter and wall thickness of 30 nm with an equiaxial composition. Magnetic characterization reveals that all the nanostructures are soft ferromagnetic, with coercivity values below 60 mT. From the angular dependence of coercivity and the relative remanence it may concluded that in both, nanowires and nanotubes the magnetization reversal mechanism undergoes a transition from one at smaller angles, involving localized nucleation by curling, and further expansion of vortex-like domain walls. At higher angles, where the applied field is almost perpendicular to the NW/NT long axis, the mechanism changes to one involving nucleation by localized coherent rotation and further expansion of transverse Bloch-like walls.

There has been an explosive growth of nanoscience and nanotechnology in the last few years, primarily because of the availability of new strategies for the synthesis of nanomaterials and new tools for their characterization and manipulation. Nanostructures and nanomaterials dominate the research scene in almost all areas of natural sciences and engineering worldwide due to their countless applications in different areas of knowledge such as medicine, pharmacology, photonic, spintronic and many others.1–3 In this sense, it is possible to see in the literature new ways to synthesize and characterize nanostructures with different morphologies and dimensionalities, and fundamentally new materials designed for specific applications.4,5

Magnetic nanowires have been widely investigated in connection with their potential applications in magnetic and electronic nanodevices.6 Magnetically hard ferromagnetic alloys containing rare earths have been specially studied in nanowire morphology for an improvement in their magnetic moment, from the angular dependence of coercivity and the relative remanence it may be concluded that in both, nanowires and nanotubes the magnetization reversal mechanism undergoes a transition from one at smaller angles, involving localized nucleation by curling, and further expansion of vortex-like domain walls. At higher angles, where the applied field is almost perpendicular to the NW/NT long axis, the mechanism changes to one involving nucleation by localized coherent rotation and further expansion of transverse Bloch-like walls.

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Although the magnetic properties of the different phases in Sm-Co alloys are well known, very few works have been reported regarding the synthesis of Sm-Co nanowires by electrodeposition (ED), particularly in aqueous medium.7 In this aspect, Cojocaru et al. have used a 1 chloride: 2 urea molar eutectic solvent mixture to carried out SmCo electrodeposition in a copper substrate and also in alumina template,8 and Chen et al. have deposited SmCo alloy at constant potential, from 1-ethyl-3-methylimidazolium chloride ionic liquid, without template.9

In order to improve the morphology and microstructure of these 1D particles, and also the magnetic and/or electronic properties, electrocrystallization conditions must be precisely determined.

In the present work, Sm-Co nanowires were fabricated by ED into AAO templates and further characterized, in order to experimentally determine the value of the reduction potential for the electrodeposition of the alloy in an aqueous medium. The effects of the electrodeposition potential applied during the synthesis process on the composition, crystallography, morphology and on the magnetic properties of the nanostructures are further investigated. This is an original result as no data related to Sm-Co electrodeposited in aqueous medium at these low potentials, and without chelating agents have been reported.

Experimental

Materials.—All reagents, SmCl3·7·H2O (Alfa Aesar), CoSO4·7·H2O (Sigma-Aldrich) and H3BO3 (Cicarello) were of analytical grade and being used without further purification. Commercial AAO porous templates (about 1010 pores cm−2), with nominal pore diameter of 200 nm, 60 μm thickness, were purchased by Whatman. Aqueous solutions were prepared by using 18.2 MΩ cm−1 resistance water (Milli-Q, Millipore, Billerica, MA), pH measurements performed with a combined glass electrode and a digital pH-meter (Altronix). Unless noted, all experiments were performed at room temperature, (300 ± 2) K (27 ± 2°C).

Nanowire/nanotube arrays are obtained by ED; a continuous layer of gold (ca. 30 nm thick) is thermally sputtered onto one side of the template to serve as an AAO/gold working electrode. ED of Sm-Co is carried out in a three-electrode cell, using aqueous solution (ED solution) of 60 mM CoSO4 + 60 mM SmCl3 in 500 mM H3BO3 as electrolyte, at pH 3 and at room temperature. A high-purity platinum plate is used as a counter electrode and a saturated Ag/AgCl...
electrode serving as the reference electrode. Deposition and cyclic voltammetry studies were performed in an Autolab302N potentiostat/galvanostat device, with the NOVA1.8 measurement software installed.

The resulting microstructures were characterized by scanning (SEM), transmission (TEM) electron microscopy, and X-ray photoelectron spectroscopy (XPS). SEM images of nanowires/nanotubes were recorded on a SEM Carl Zeiss Sigma, and the atomic percentage of Sm and Co determined by energy-dispersive X-ray spectroscopy (EDS). TEM observations, images and electron diffraction patterns were performed in a Philips CM 200 UT microscope.

XPS experiments were carried out using a K-Alpha X-ray photoelectron spectroscope (Thermo-Fisher Scientific Co.), with a monochromatized Al-Kα X-ray source in vacuum of 10⁻⁸ mBar.

Room temperature magnetic hysteresis loops were measured in a vibrating sample magnetometer (VSM) Lake-Shore 7300, with a maximum field up to 1.5 T, to determine the coercive field $H_C$ and the relative magnetization or squareness $MS/MB$ of Sm$^{3+}$ and Co$^{2+}$.

Results and Discussion

Mainly two factors are found to affect the deposit composition: one of them is the reduction potential value of the species involved (Sm, Co) and the other one is the concentration ratio of the components in the feeding solution used for the electrodeposition. The reduction potentials are very different ($E_{Sm^{3+}/Sm}$ = −2.30417 and $E_{Co^{2+}/Co}$ = −0.28018) making difficult to obtain the alloy. An equimolar concentration of Sm:Co is used in the ED solution as indicated by Mishra et al. These authors conclude that at lower Co concentration (relative to the rare earth), the process rate decreases because there is not enough Co to catalyze the rare earth (RE) deposition.

Brenner describes the condition where the less noble element is preferentially deposited, and defines this process as anomalous deposition which is a type of abnormal deposition. The induced co-deposition represents another type of abnormal deposition in which an element that cannot be deposited in its pure form, is co-deposited as an alloy.

Figure 1 shows the cyclic voltamperograms for the blank solution (dotted line) and for the bath containing Sm$^{3+}$ and Co$^{2+}$ ions, measured at 50 mV·s⁻¹ (dashed line). A three electrode electrochemical cell was used with a steel (1 cm²) working electrode, a high-purity platinum plate as a counter electrode and saturated Ag/AgCl as the reference electrode. As can be seen, blank solution produces an almost null trace, while for the ED solution three reduction peaks at −1.0 V, −1.3 V and −1.5 V can be observed, as is clearly shown in the insert.

The peak located at −1.0 V is related to Co$^{2+}$ to Co reduction. The peaks observed at −1.3 V and −1.5 V arise from the reduction of Sm$^{3+}$ and Sm$^{2+}$. There is evidence that the Sm-Co deposition follows an induced co-deposition behavior, which was also reported in RE-transition metal alloys deposition such as Co with Tb$^5$ or Gd$^6$, where cobalt deposition is inhibited once the rare earth deposition starts. RE electrodeposition at potentials which are more positive than the equilibrium reduction potential is enabled by the catalytic action of cobalt, similar to the induced co-deposition mechanism observed in molybdenum and tungsten. Liu et al. observed the deposition of SmCo and NiFe at lower potential with respect to the reduction potential of Sm(III) and Nd(III), which is in the range of −2.5 V vs. Saturated Calomel Electrode (SCE). As a consequence, they proposed that a mixed intermediate metal is formed. This is why the peaks observed at approximately −1.3 V and −1.5 V are attributed to the reduction of Sm$^{3+}$ to the intermediate hydroxide species SmCoOH (Sm$^{2+}$) and the reduction of this intermediate compound to Sm$_{\text{red}}$, respectively. This idea is supported by the absence of a buffer solution and the concomitant increment in the local OH⁻ concentration, as a result of the hydrogen electrochemical reaction. It should be noted that although boric acid is used as electrolyte support in the electrodeposition, the pH of the solution is around 3, far away from the pKa value of boric acid, pKa = 9.24, so it is not a buffer solution. In the present work the solution is not stirred because Panzeri et al. observed that stirring promotes a high depletion in the Sm content. Additionally, Kolb et al. attribute the deposition of some metals at more positive potential values to the differences in the work function value ($\phi_M$) between the metallic electrode and the metallic monolayer deposition. The difference $\phi_{Sm}-\phi_{Co} = 2.4$ eV implies a potential shift of $\Delta E_{\text{red}} = 1.2$ V ($\Delta E_{\text{red}} = 0.5$ V (eV)⁻¹ $\phi_M$) to more positive values, that is, from $E_{\text{red,Sm}} = −2.3$ V to $−1.1$ V. This shift is explained by the generation of a partial charge in the adatoms, giving a ionic contribution to the chemical bond which reinforces the induced co-deposition effect.

Based on these results five potential values were selected to produce nanostructures by elec-trodeposition. These values (−0.8 V, −0.9 V, −1.0 V, −2.0 V and −3.0 V) are marked in Figure 1.

Figure 2 shows typical $J$ vs. $t$ transient curves recorded in the course of the potentiostatic alloy deposition at −0.9 V and −3.0 V. The electrodeposition processes exhibit characteristic stages: first, a maximum of cathodic current corresponding to the initial stage is observed, followed by a stage of non-monotonic current decrease, eventually reaching a constant current stationary regime, which corresponds to the nanostructure growth inside the cylindrical AAO channels.

At a potential of −3.0 V, variations in the current are observed due to the H₂ discharge. All electrodeposition experiments were stopped when the total charge value was 800 mC; and as a consequence, the total electrodeposition times are different and depend on the applied potential, as illustrated in the inset.
Morphology and microstructure.—Figure 3 shows SEM micrographs corresponding to the different electrodeposited SmCo arrays, after the partial dissolution of the alumina membrane in NaOH 1 M during 60 min. The arrays consist of ordered, parallel and densely distributed nanowires for $-0.9$ V (Figure 3a) and nanotubes for $-3$ V (Figure 3b), indicating that pores were optimally filled. Details about the arrays size and morphology are listed in Table I.

Nanowires obtained at $-0.8$ V are short and quite irregular. Nevertheless, the resulting nanowires are long and perfectly ordered for a potential of $-0.9$ V, as can be observed in Figure 3a. At this potential Sm-Co nanowires are abundant, uniform and densely packed in a large scale.

At a potential value of $-1.0$ V long, perfectly ordered nanowires are obtained mixed with also regular nanotubes. There are no previous reports on Sm-Co 1D nanostructure obtained at such low potentials.

In this condition, the background of hydrogen discharge associated to high potential values ($-2$ V or $-3$ V) can be avoided without the use of expensive and impractical procedures.

When the potential applied is $-2.0$ V or $-3.0$ V the resulting nanostructures are nanotubes 195 nm outer diameter and 30 nm wall thick, leading to an internal diameter of 135 nm and to a factor $\beta = (D_{int}/D) = 0.7$ (Figure 3b). This factor determines the magnetization reversal mechanism in the NT as will be seen below. This NT morphology, appearing for high applied potentials promoting rapid nanostructure growth, probably arises from the strong hydrogen evolution. Fukunaka et al. and Davis et al. observed in electrodeposition of transition metal nanotubes, that the hydrogen evolution played a major role in determining wire or tube formation.

The nominal composition, obtained by EDS, is plotted as a function of the applied potential in the electrodeposition process and is shown in Figure 4. At electrodeposition potentials as low as $-0.8$ V, $-0.9$ V and $-1.0$ V, the nanostructure nominal composition approximately corresponds to Sm 10 at.\% and Co 90 at.\%, in agreement with that of the Sm$_2$Co$_7$ equilibrium phase in this binary system. On the other hand, at potentials usually applied for rare earth deposition such as $-2.0$ V and $-3.0$ V, the nominal composition changes to the equiatomic one, Sm 50 at.\% and Co 50 at.\%.

Table I. Morphological parameters corresponding to the electrodepositions performed at different potentials. The cylinder length $L$, the internal structure (nanowires NWs or nanotubes NTs), their external diameter $D$, the NTs wall thickness, the respective aspect ratios $a_R$ (= $L/D$) and $\beta$ (= $(D - 2t_W)/D$) factors are listed.

| $E$ [V] | $L$ [$\mu$m] | Morphology | $D$ [nm] | $t_W$ [nm] | $a_R$ | $\beta$ |
|---|---|---|---|---|---|---|
| 0.8 | 1.0 ± 0.5 | NW | 178 ± 10 | - | 5.5 | - |
| 0.9 | 6 ± 1 | NW | 180 ± 10 | - | 33 | - |
| 1.0 | 5 ± 1 | NW/NT | 160 ± 10 | 35 ± 3 | 33 | - |
| 2.0 | 20 ± 3 | NT | 190 ± 10 | 30 ± 3 | 100 | 0.7 |
| 3.0 | 18 ± 3 | NT | 205 ± 10 | 31 ± 3 | 100 | 0.7 |

Figure 4. EDS results of atomic percent of Co expressed as a function of applied potential.

There is some evidence that the deposition rate of Sm in Sm-Co alloys considerably increases when the potential applied is near to $-2.0$ V. In this sense, our results are consistent with previously published data for electrodeposition of Sm-Co films from aqueous media.

In order to determine the degree of crystallinity and the phases in the nanostructures, XRD experiments were performed for all the samples. Diffractograms (not shown) were practically featureless, with only a few small peaks arising from the alumina template or the gold layer.

Then, crystallographic properties were further explored by TEM. Figure 5 shows bright field images of typical nanostructures resulting for a potential value of $-0.9$ V. Figure 5a and Figure 5b. A subtle film is frequently found around the nanostructures, which contains CoO, most probably formed during the sample dissolution of the alumina membrane in NaOH 1 M.
Figure 6. (a) Dark field image of a NT obtained at $-3 \, \text{V}$ taken with a spot belonging to the crystalline nanophase ($\sim 5 \, \text{nm}$) indicating that these small crystallites are CoO formed onto the NTs and NWs surfaces. (b) Electron diffraction rings corresponding to the CoO phase (arrows).

Figure 7 shows XPS experiments for nanowires in the region of Sm, Figure 7a, and Co signals, Figure 7b, before and after stripping the surface of the nanowires with an Ar beam. Sm peaks before stripping, which correspond to $3d_{3/2}$ and $3d_{5/2}$, appear at 1084.3 eV and 1111.5 eV ($\Delta E = 27.2$ eV), respectively, and at 1084.0 eV and 1110.8 eV ($\Delta E = 26.8$ eV) after that procedure. The slight change and the decrease in the peaks separation, are related to the fact that the metallic character of the surface increases as a result of the stripping.32–34

These energy values are slightly higher than expected for metallic Sm, because in the SmCo alloy, Sm is attached to Co, which is more electronegative. In addition, as shown in Figure 7b, the peaks corresponding to cobalt oxide were observed, and with the first pickling, the Co metallic signal increases considerably.35,36 It should be noted that as the AAO template is dissolved with NaOH 1M in the present protocol, then, the metallic oxides could be formed at this stage.

Magnetic hysteresis properties.—The effect of the different potentials applied during electrodeposition on the magnetic properties was investigated by measuring the room temperature hysteresis loops. Low absolute potential values lead to higher Co content in the amorphous phase and also to NW morphology, both features promoting higher coercivity values. Figure 8 shows typical hysteresis loops of ordered Sm-Co nanowires, measured inside the AAO templates, for two extreme applied potentials as 0.9 V (a) and $-3.0 \, \text{V}$ (b). The hysteresis loops are measured with the external field applied parallel (PA) and perpendicular (PE) to the major axis of nanowires/nanotubes.

Considering that NWs and NTs are amorphous quite low crystalline anisotropy may be expected and only shape anisotropy is considered. Shape anisotropy and the corresponding shape anisotropy field in NWs...
are estimated as:

\[ K_{\text{sh}}^{\text{NW}} = \frac{1}{2} \Delta N \mu_0 M_S^2 \]  

[1]

\[ \mu_0 H_{\text{sh}}(T) = \frac{1}{2} \Delta N \mu_0 M_S(T) \]  

[2]

with \( \Delta N \) the difference between of demagnetizing factor along the major wire axis and that in a normal direction, and \( \mu_0 M_S \) the NWs’ saturation polarization. For NWs with nominal composition close to the Sm2Co17 phase and an aspect ratio of 33, the saturation polarization may be approximated by the value corresponding to this phase \( \mu_0 M_{\text{SNW}} = 1.25 \) T (298 K). The desmagnetizing factor of finite nanowires along the major length may be estimated as proposed by Landeros et al., leading to

\[ N = 0.044. \]

Then, \( \Delta N^{\text{NW}} = 0.43, K_{\text{sh}}^{\text{NW}} = 2.67 \times 10^7 \) J m\(^{-3}\) and \( \mu_0 H_{\text{sh}}^{\text{NW}}(T) = 0.268 \) T.

For nanotubes obtained at \(-2.0\) V and \(-3.0\) V the demagnetizing factor along the major axis is negligible (\( \Delta N^{\text{NT}} = 0.5 \)) and the shape anisotropy constant and field are respectively \( K_{\text{sh}}^{\text{NT}} = 0.95 \times 10^7 \) J m\(^{-3}\) and \( \mu_0 H_{\text{sh}}^{\text{NT}}(T) = 0.17 \) T.

These magnitudes indicate that both nanostructures are quite soft, as expected for amorphous samples, and that larger coercive fields and magnetostatic shape effects are expected in nanowires than in nanotubes, as observed in Figure 8. In both types of nanostructure the effective easy axis is parallel to the long axis.

Figure 9 summarizes coercivity (\( \mu_0 H_C \)) and squareness (\( S = M_r/M_s \)) for SmCo nanotube and nanowires arrays, as functions of the electrodeposition potential. As observed in Figure 8 for the extreme potentials values, no significant difference between the coercivity in configurations PA and PE is observed in all the cases. Coercivity decreases as the potential applied in the electrodeposition increases due to the transition from NW to NT morphology and the decrease in the Co content in the nanostructure. Squareness (S) also decreases for higher potential while the difference between the two orientations becomes smaller as discussed in Figure 8.

The magnetization reversal mechanisms operating in these amorphous 1D nanostructures, with NW and NT morphologies and also different compositions, may be characterized by measuring the coercive field and the relative remanence (S) as functions of the angle \( \theta \) between the long axis of NWs and NTs, and the applied magnetic field. The results for samples synthesized at different electrodeposition potentials are shown in Figure 10. It may be observed that, despite the different nanostructures’ morphology and composition, the shape of curves in Figure 10a \( \mu_0 H_C(\theta) \) are quite similar, and the same is found for curves S(\( \theta \)) shown in Figure 10b.

Coercivity angular variation shown in Figure 10a is similar to curves reported by other authors in NWs and NTs. Curves in Figure 10b, displaying the monotonic decrease of the relative remanence with the angle \( \theta \), are also similar to those reported by these latter authors.

The non-monotonic behavior of curves \( \mu_0 H_C(\theta) \) arises from the transition from an initial reversal regime to another one, taking place at a given critical angle depending on the sample geometry and magnetic hardness. The initial reversal mechanism involves the localized nucleation by curling, and further expansion of vortex-like domain walls. At higher angles, where the applied field is almost perpendicular to the NW/NT long axis, the mechanism changes to one involving nucleation by localized coherent rotation and further expansion of transverse Bloch-like walls. This transition is observed in NWs...
and NTs at somewhat different critical angles $\theta_{\text{CT}} \approx 70\text{–}75^\circ$. In the case of magnetically soft NTs, this critical angle increases with the outer diameter, being about $55^\circ$ for a diameter of $d = 50$ nm while for $d = 65$ nm, the critical transition angle reaches $\theta_{\text{CT}} = 67^\circ$. One should note that the outer and inner diameters of a NT array will highly influence the type of DW that is energetically more favorable to nucleate. In addition to the transition in the reversal mechanism, another factor promoting this type of non-monotonic coercivity curves is the dipolar interactions between nanocylinders, as reported by Bender et al., especially in our case of large diameter 1D nanostructure.

**Conclusions**

This work demonstrates for the first time that it is possible to synthesize Sm-Co 1D nanostructures by electrodeposition of the alloy in aqueous medium, at quite low potential values and without chelating agents.

For the first time long, uniform and densely packed Sm-Co nanowire arrays were prepared by electrodeposition inside the cylindrical pores of AAO templates, at quite low electrodeposition potentials such as $-0.8 \text{ V}$, $-0.9 \text{ V}$ and $-1.0 \text{ V}$. Samples were amorphous NWs, with a nominal composition close to that of the equilibrium stoichiometric phase Sm$_2$Co$_{17}$. At very high potential as $-3.0 \text{ V}$, long and regular nanotubes $190 \text{ nm}$ in outer diameter and of $30 \text{ nm}$ thick walls form with the equiatomic SmCo composition, mainly due to the strong hydrogen evolution.

NWs and NTs are soft ferromagnetic, with the easy axis parallel to the sample major axis and magnetic hardness is governed by the shape anisotropy through the saturation polarization (Co content) with little influence of geometry (aspect ratio).

The angular variation of the coercive field with the angle $\theta$ between the major axis of the nanostructure (easy axis direction) and the applied magnetic field.

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**Figure 10.** Angular dependence of (a) the coercive field and (b) the relative remanence, corresponding to the NWs and NTs electrodeposited, $\theta$ is the angle between the major axis of the nanostructure (easy axis direction) and the applied magnetic field.
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