Electrodeposition of Magnetic SmCo Films from Deep Eutectic Solvents and Choline Chloride-Ethylene Glycol Mixtures

Gabriele Panzeri,a Matteo Tresoldi,a Christian Rinaldi,b and Luca Magagnin a,b,*

a Dipartimento di Chimica, Materiali e Ingegneria Chimica Giulio Natta, Politecnico di Milano, 20131 Milan, Italy
b Dipartimento di Fisica, Politecnico di Milano, 20133 Milan, Italy

Suitability of mixtures based on ethylene glycol and choline chloride is evaluated for the electrodeposition of magnetic SmCo alloys. Deep eutectic solvents (DES) are characterized by a wide electrochemical window, allowing the electrodeposition of metallic elements having a highly negative reduction potential. Electrodeposition of unconventional metals with reduction potential well below hydrogen evolution one is, in principle, feasible (e.g. Sm). Moreover, the limited presence of water during the growth of SmCo alloys may reduce oxidation and consequently improve the magnetic properties with respect to the employment of an aqueous bath. Deposition composition is strongly affected by process parameters and bath composition. Increasing choline chloride concentration and/or adding glycine to the solution allows to achieve higher Sm content in the deposit. Potentiostatic electrodeposition has been carried out in a conventional three electrodes cell, with potential interval selected from $-0.7 \text{ V} \text{ vs Ag wire}$ to $-0.95 \text{ V} \text{ vs Ag wire}$, keeping the bath under stirring at 70$^\circ C$. Magnetic SmCo films with composition from 0 to 50 wt% Sm have been obtained with coercivity up to 270 Oe. 

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The role of non-aqueous electrolytes is gaining importance due to their ability to overcome limitations normally observed in the electrodeposition processes from aqueous media. In particular, this is true for the metal elements having reduction potential well below hydrogen evolution one. Ionic liquids are in fact characterized by wider electrochemical windows limiting the onset of secondary processes suppressing the metal reduction one.1–3 One practical example is the electrodeposition of rare earth metals and their alloys (e.g. SmCo). Gomez et al. and Cojocaru et al. proposed a study on the electrodeposition of SmCo film from a deep eutectic solvent (DES) based on choline chloride and urea (1ChCl:2U).4–6 The suitability of such a system for the electrodeposition of both single metals (Sm and Co) and alloy (SmCo) was verified. Depositions were characterized by a high Sm content, i.e. 46–79 wt% Sm, while the optimal composition for the best magnetic performances is 23–25 wt% Sm, corresponding to the Sm$_2$Co$_{17}$ phase.7–9 In literature, works on this alloy from both water-based10–16 and non-aqueous solution are reported.17 In the former case, fundamental is the employment of glycine as complexing agent as shown by Wei et al.15–16 However, in-plane coercivities in the order of $H_{c//} = 100$ Oe were observed with the formation of SmO, Sm(OH)$_3$ and Co(OH)$_2$.14 These values are referred to films produced by electrodeposition, an interesting technique from the fabrication point of view but with a huge limitations in terms of compositional and purity control with respect to traditional methods as casting or sputtering through which hard SmCo magnets are obtained ($H_{c//} \sim kOe$).

This work consists in the study of different mixtures based on ethylene glycol and choline chloride (1ChCl:2EG and 1ChCl:4.5EG) in order to evaluate the effect of ChCl concentration on SmCo codeposition and its magnetic properties. Studies on the physical properties of solutions with different ChCl:HBD (Hydrogen Bond Donor) molar ratio are reported in literature. It has been shown that reducing the amount of choline chloride from 33% (1ChCl:2EG) to 15% (1ChCl:5EG) molar, viscosity is reduced from 36 cp to 15 cp at 20$^\circ$C with a slight increase in the conductivity, from 7.61 mS cm$^{-1}$ to 8.14 mS cm$^{-1}$.18,19 Thus, by changing the choline chloride concentration in the interval considered, the physical properties of the system are maintained, considering Eg as HBD. The different concentration of hydrogen bond acceptor (HBA) allows to have an additional control on alloy film composition. Moreover, the employment of a lower quantity of choline chloride, highly hygroscopic, reduces water contamination of the solution. Eventually, glycine is added to evaluate its effect in a system different from an aqueous one.

Experimental

Electrochemical characterization of the bath and deposition experiments have been carried out using AMEL2553 potentiostat/galvanostat with VAPeak software. Experimental apparatus was a conventional three electrodes electrochemical cell, whereas for the preliminary study on the deposit composition dependence on bath agitation, a rotating electrode was employed. For cyclic voltammetry CV, a platinum wire was employed as both anode and cathode. Deposition of magnetic films were performed on copper substrate using a conductive mixed metal oxide mesh (TiO$_2$/RuO$_2$/IrO$_2$) as counter electrode. Silver pseudo-reference electrode was employed for all the experiments, potential values were referred to Ag wire. Prior each experiment, all electrodes were cleaned with a 10 wt% HNO$_3$ aqueous solution. Copper substrates have been degreased and subsequently etched in 15 wt% H$_2$SO$_4$, 3 wt% H$_2$O$_2$ aqueous solution. Two different solvents were considered, respectively 1ChCl:4.5EG and 1ChCl:2EG molar ratio. Solutions were prepared under constant stirring at 70$^\circ$C, 0.04 M cobalt(II) chloride hexahydrate [CoCl$_2 \cdot 6$H$_2$O] and 0.04 M samarium(III) chloride hexahydrate [SmCl$_3 \cdot 6$H$_2$O] were added subsequently. Glycine [C$_2$H$_5$NO$_2$] (0.12 M) was added to evaluate its effect on deposition process and film properties. All reagents employed were of analytical grade. Cyclic voltammograms were carried out at 10 mV s$^{-1}$ and different deposition potentials were considered for the potentiostatic deposition in the range from $-0.7 \text{ V} \text{ vs Ag wire}$ to $-0.95 \text{ V} \text{ vs Ag wire}$. For all experiments, working temperature was kept at 70$^\circ$C. Composition was measured by X-Ray Fluorescence (XRF) with a Fischerscope X-ray XAN. Film surface morphology and crystal structure have been investigated by means of Scanning Electron Microscopy (SEM) and X-Ray Diffraction (XRD) analysis (Philips model PW1830. Cu-Ka1 = 1.54058 Å). Magnetic properties have been evaluated using a Vibrating Sample Magnetometer (VSM, model EZ9 from MicroSense LLC, Massachusetts, USA).

Results and Discussion

Cyclic voltammetry was performed in solutions containing individual metal salts; system suitability was thus evaluated for the electrodeposition of Sm and Co films. 1ChCl:2EG DES with no metal salts have been taken as reference to evaluate the onset of reduction peaks in the pure metal solution. For blank solutions, a typical CV curve was obtained ($H_{c//} \sim 100$ Oe), resembling the behavior of a conductive mixed metal oxide mesh (TiO$_2$/RuO$_2$/IrO$_2$) as counter electrode. Silver pseudo-reference electrode was employed for all the experiments, potential values were referred to Ag wire. Prior each experiment, all electrodes were cleaned with a 10 wt% HNO$_3$ aqueous solution. Copper substrates have been degreased and subsequently etched in 15 wt% H$_2$SO$_4$, 3 wt% H$_2$O$_2$ aqueous solution. Two different solvents were considered, respectively 1ChCl:4.5EG and 1ChCl:2EG molar ratio. Solutions were prepared under constant stirring at 70$^\circ$C, 0.04 M cobalt(II) chloride hexahydrate [CoCl$_2 \cdot 6$H$_2$O] and 0.04 M samarium(III) chloride hexahydrate [SmCl$_3 \cdot 6$H$_2$O] were added subsequently. Glycine [C$_2$H$_5$NO$_2$] (0.12 M) was added to evaluate its effect on deposition process and film properties. All reagents employed were of analytical grade. Cyclic voltammograms were carried out at 10 mV s$^{-1}$ and different deposition potentials were considered for the potentiostatic deposition in the range from $-0.7 \text{ V} \text{ vs Ag wire}$ to $-0.95 \text{ V} \text{ vs Ag wire}$. For all experiments, working temperature was kept at 70$^\circ$C. Composition was measured by X-Ray Fluorescence (XRF) with a Fischerscope X-ray XAN. Film surface morphology and crystal structure have been investigated by means of Scanning Electron Microscopy (SEM) and X-Ray Diffraction (XRD) analysis (Philips model PW1830. Cu-Ka1 = 1.54058 Å). Magnetic properties have been evaluated using a Vibrating Sample Magnetometer (VSM, model EZ9 from MicroSense LLC, Massachusetts, USA).
ChCl:EG system reported in Figure 1 was observed: the lower potential limit is set to −1.5 V vs Ag wire because of the significant noise in the signal for more negative potential, related to hydrogen evolution. Results match with the data reported by Badea et al.,\textsuperscript{20} where a lower amount of ChCl in solution corresponds to a wider electrochemical window. Adding single metal salts, there were no significant variations in the CV. For 0.04 M cobalt(II) solution, no reduction peak was observed. On the other hand, there was a significant anodic peak assessed to Co film stripping for the 1ChCl:4.5EG solution, while it was less intense for 1ChCl:2EG one. Considering the addition of 0.04 M samarium(III), no reduction peaks and corresponding anodic peaks were observed for both the solutions. Same results were observed for higher salt concentrations, e.g. 0.08 M samarium(III). Thus, such a system seems to be not suitable for the deposition of metallic samarium film on Pt in the considered potential range. More negative cathodic limits were selected but differences in the electrochemical behavior were not observed.

SmCo codeposition was firstly studied from a solution containing 0.04 M SmCl\(_3\)·6H\(_2\)O and 0.04 M CoCl\(_2\)·6H\(_2\)O. Voltammogram curve showed a wide reduction peak at −0.9 V vs Ag wire for the 1ChCl:4.5EG solution. Even though it is not possible to identify two different reduction peaks, respectively corresponding to samarium and cobalt (Figure 2), the presence of both Sm and Co ions helps the codeposition of SmCo through an induced codeposition mechanism, as observed for SmCo alloys from molten salts.\textsuperscript{21} As previously shown, the 1ChCl:2EG curve is shifted toward positive potential values showing a reduction peak with lower intensity at −0.6 V vs Ag wire. In both cases, during the forward scan (from 0 to −1.5 V vs Ag wire), a film was formed on Cu substrate and subsequently removed rising the potential toward +0.5 V vs Ag wire.

Subsequently, 0.12 M glycine has been added to both solutions. Different studies on aqueous solutions report the 3[Gly]:[Co]\textsuperscript{2+} ratio as the most suitable for codeposition of Co and Sm; a higher concentration of glycine would in fact hinder the Sm deposition.\textsuperscript{13–15} The same concentration ratio has been selected also for our systems despite based on different chemistry. From the CV curve (Figure 3), it is evident the effect of glycine on the electrochemical behavior of the bath, especially for the 1ChCl:4.5EG solution. Voltammogram curve presents two reduction peaks at −0.85 V vs Ag wire and −1 V vs Ag wire, attributed to Co and Sm reduction respectively. Moreover, curves shift toward positive potential, favoring the Sm codeposition in the deposit. In the 1ChCl:2EG curve, a first reduction peak is found at −0.75 V vs Ag wire and a smaller, less intense, at −0.92 V vs Ag wire. With respect to glycine-free solution, reduction process is favored and a more intense anodic peak is observed.

Glycine effect on deposit composition was evaluated by means of potentiostatic deposition in the potential window from −0.7 V vs Ag wire to −0.95 V vs Ag wire. Plating process was carried out under stirring, stagnant conditions were found not suitable for SmCo electrodeposition. Similar results were observed for relatively high negative potential with formation of a black oxide powder at −0.9 V vs Ag wire. Consequently, deposits were inhomogeneous in composition and thickness; this may be probably due to the relatively high viscosity of ChCl:EG system. On the other hand, because of the hygroscopic nature of choline chloride, a certain amount of water is always present in solution and hydrogen evolution at the cathodic surface can occur during the electrodeposition process affecting surface morphology. For all these reasons, stirring plays a fundamental role as shown from the preliminary study on 1ChCl:4.5EG solution where increasing stirring (rotating disk electrode: from 10 to 200 RPM) decreases the Sm concentration in the deposits.\textsuperscript{22} After having defined the solution agitation effect, potentiostatic depositions were carried out in a conventional three-electrode electrochemical cell where the magnetic stirrer was set to constant value (300 RPM), different deposition potentials were selected from −0.7 V vs Ag wire to −0.95 V vs Ag wire. Four different solutions were considered to study the role of
choline chloride concentration and the presence of glycine in the bath. Sm concentration and deposition potential match with the voltammetry curve: increasing the driving force of the plating process leads to higher samarium content in the deposit. Film composition was also highly dependent on the selected system (Table I). The majority of the deposits showed a metallic aspect although the film color was blackish for high Sm content; this may be an indication of oxygen presence in the film. On the other hand, by increasing further the deposition potential (−0.9/−0.95 V vs Ag wire), the formation of oxide/hydroxide powders on the cathodic surface was observed. Finally, for three of the four solutions it was possible to obtained the target composition of 20–25 wt% Sm. In this view, ethylene glycol represents a better alternative to urea considering choline chloride as metal halide for Co–rich SmCo alloys. Gomez et al. and Cojocaru et al. showed a study on 1ChCl:2U solution where the minimum samarium concentration obtained was 45 wt% Sm, whereas Sm2Co17 is around 23 wt% Sm.4,5 Moreover, the employment of ethylene glycol instead of urea lowers significantly the viscosity thus increasing the conductivity.

Both choline chloride concentration and glycine affect significantly film composition. As observed from the cyclic voltammograms, where the curve is shifted toward more positive values, higher samarium content can be obtained for a given deposition potential. The higher ChCl concentration seems to bring to a different complexation between Sm and Co ions. For 1ChCl:4.5EG, its concentration is in fact limited to a maximum of 2 wt% Sm; for potential values lower than −0.9 V vs Ag wire, oxide powder was formed. On the other hand, it was possible to reach 50 wt% Sm increasing ChCl to 33% molar (1ChCl:2EG): this solvent composition corresponds to the eutectic point and it is comparable with the 1ChCl:2U solution. The amount of glycine added was three times the cobalt concentration; such a ratio was reported as effective for the codeposition of SmCo in aqueous system. Wei et al.13–15 demonstrated how a higher glycine quantity leads to a lower Sm content in the deposit; the same [Gly]/[Co]2+ has been adapted to our system. Results in potentiostatic conditions are in agreement with cyclic voltammetry where a more evident reduction process was observed. At a given deposition potential, Sm content is significantly increased; this is particularly true for the 1ChCl:4.5EG solution. Moreover, film growth is favored, thicknesses in the order of micrometers can be easily achieved while for the glycine-free solution the maximum thickness obtained is around 1 μm, especially for relatively high Sm content film (Sm > 15 wt%). The presence of glycine thus affects the growth kinetic; changes in the morphology are consequently expected. SEM images of deposits having 20 wt% Sm, obtained from two different solutions, are shown. Glycine addition resulted in a more compact and smooth surface showing bigger cracks but less in number. Without additive, film morphology is granular with a more disordered structure where a higher amount of cracks, smaller in size, are observed (Figure 4).

Table I. Range of film composition in the potential interval −0.7 V vs Ag wire to −0.95 V vs Ag wire, at 70°C and stirred conditions: 0.04 M CoCl2 · 6H2O, 0.04 M SmCl3 · 6H2O, 0.12 M glycine.

| Solution     | wt% Sm |
|--------------|--------|
| 1ChCl:4.5EG  | 0.5–2  |
| 1ChCl:4.5EG (Gly) | 5–29  |
| 1ChCl:2EG    | 7–25   |
| 1ChCl:2EG (Gly) | 18–44 |

The summary of samples and their in-plane magnetic properties. Pure Co deposit from glycine bath was measured as reference. As shown in Figure 6a, the film presents a magnetic hard axis in the out-of-plane direction, while the in-plane hysteresis loop shows a coercivity of 73 Oe (5.8 A/m). The S-shaped hysteresis loop (relatively low squareness) could be ascribed to the polycrystalline nature of the electrodeposited Co film, as confirmed by the XRD spectrum of Co (Figure 5). In general and no matter the solution used for the growth, the addition of the rare-earth element Sm improves the in-plane coercivity of the deposits, as shown in Figures 6b–6c for the SmCo samples. As in Co, the out-of-plane direction results to be a hard axis for the magnetization (Figures 6b–6c). The SmCo deposited in the glycine-containing solution shows a coercive field of 100 Oe (8.0 kA/m), higher than Co film and compatible with those reported in literature for electrodeposition from aqueous baths.13–15 On the other hand, SmCo film grown in a 1ChCl:2EG solution without glycine presents a significantly higher coercivity, while the SmCo obtained from aqueous baths shows coercivities well below 50 Oe.6–8

Figure 4. Deposit morphology of SmCo alloys (20 wt% Sm) obtained from 1ChCl:2EG solution. a) Glycine-free solution. b) Glycine-containing solution.

Figure 5. XRD spectra of Co electrodeposited from glycine containing bath. SmCo electrodeposit with 20 wt% Sm from glycine-free bath and SmCo electrodeposit with 20 wt% Sm from glycine-containing bath in 1ChCl:2EG.
Table II. Magnetic properties obtained by the in-plane hysteresis loops of the electrodeposited films.

| Sample       | Solution   | wt% Sm | in-plane $H_c$ [Oe] | Loop squareness $M_r/M_{sat}$ |
|--------------|------------|--------|---------------------|-------------------------------|
| Co           | 1ChCl:2EG (Gly) | 0      | 73                  | 0.25                          |
| SmCo (gly-free) | 1ChCl:2EG   | 20     | 270                 | 0.21                          |
| SmCo (gly)   | 1ChCl:2EG (Gly) | 20     | 100                 | 0.27                          |

Figure 7. Comparison between the hysteresis loops of the three electrodeposited films.


coercive field $H_c = 270$ Oe (21.5 A/m), in agreement with the value reported for the 1ChCl:2U system. The out-of-plane direction is again a hard axis for the magnetization (Figure 6b). A comparison between the in-plane M(H) curves for the three samples is shown in Figure 7. The different magnetic properties of the SmCo films can be related to the extremely different morphology of the two, as shown in Figures 4a–4b. While the sample with glycine has smoother surface and presents a higher crystalline ordering, the film grown without glycine is granular and disordered. The higher concentration of defects in the latter could be responsible for a significant pinning of the magnetization and thus of the increase of the coercivity. The misalignment of grains could also be the cause for the reduction of the remanent magnetization and thus of the increase of the coercivity. The misalignment of grains could also be the cause for the reduction of the remanent magnetization and thus of the increase of the coercivity.

Conclusions

Choline chloride/ethylene glycol based solutions have been proven to be suitable for SmCo codeposition: films with Sm content corresponding to Sm$_2$Co$_{17}$ composition have been obtained. The addition of choline chloride and glycine promotes the SmCo codeposition, with an increase in Sm content in the films. Glycine promotes also film growth by increasing the maximum achievable thickness: the resulting film shows a smooth morphology, even if cracks are present. From the point of view of magnetic properties, SmCo films are ferromagnetic. The addition of the rare-earth element Sm to Co increases the in-plane coercive field with coercivity values up to 270 Oe.

ORCID

Luca Magagnin https://orcid.org/0000-0001-5553-6441

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