Synthesis and characterization of novel electrodeposited catalytic materials for hydrogen evolution reaction

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Novel electrodeposited materials as NiMoW, NiCoMnB, CoMnVB, CoVB, CoNiMoW and NiCoMoW are reviewed. The electrolytes and electrolytic conditions for obtaining of these deposits were presented. Morphology and component distribution in the composites was observed by SEM and EDX analyses. The electrocatalytic activity for HER in different medium was estimated by polarization curves. The best electrocatalytic results for HER were shown by CoNiMoW and NiMoW systems.

Key words: Ni- and Co-based electrodeposits, multilayers, composite, hydrogen evolution.

INTRODUCTION

There is a huge number of investigations on the hydrogen evolution reaction (HER) due to its technological importance. The activity of the cathodic material possessing low hydrogen evolution overpotentials is one of the most important characteristic of the electrodes. Ni and Ni-based alloys have been extensively studied for HER [1-31] due to their good electrocatalytic properties and corrosion stability in alkaline solutions. It is proposed that the high catalytic activity is caused by an increased amount of amorphous phase surrounding the Ni crystals [6, 9], i.e. the crystal size is correlated to the catalytic activity of hydrogen evolution in alkaline solutions. The presence of TiO₂ particles in amorphous Ni-P layers [11, 14] and incorporated Ti particles in Ni deposit [13] lead to an increase of the hydrogen evolution rate in comparison with conventional Ni and Ni-P deposits. Also, the increase in activity occurring after heating of Ni-P + TiO_2 + Ti layers is related to TiO₂ reduction and formation of nonstoichiometric Ti oxides [15]. In order to improve the catalytic properties of nickel electrodes, W [16, 17] and Mo [18-27] are co-deposited with Ni. Other materials characterized by better catalytic behaviour are Co and Ni-Co based composites [28-30]. Both composites and alloys of transition metals based on Ni and Co possess higher electrocatalytic activity for HER than pure metals. This phenomenon corresponds not only to the increase of surface roughness but to the electronic structure of materials. The electron transfer occurs from the less

The aim of this work is to make a review of own novel electrosynthesized Ni- and Co-based materials suitable as catalysts for HER.

ELECTRODEPOSITION OF Ni BASED SYSTEMS

Codeposited TiO_x particles

The interest in electrodeposition of tungsten with nickel is increasing due to the technological properties of the obtained materials such as hardness [32, 33], corrosion resistance [34, 35], activity as hydrogen electrodes [16, 17], etc. The initial treatment of the substrate strongly influences the deposit characteristics. For example, thin electrodeposited Ni-W films on nanomodified carbon fibre showed more developed surface and better electrocatalytic activity compared to that of pristine carbon fibres [16]. As it was mentioned above, composite layers obtained by electrolytic codeposition of nickel and titanium powder from an electrolyte containing Ti powder suspension [36, 37] exhibited higher electrochemical activity for HER [13–15] compared with that of nickel coatings. Their high activity in this process was attributed to the developed electrode surface due to incorporation of Ti powder into the nickel matrix as well as to the presence of nonstoichiometric titanium oxides and intermetallic Ni-Ti compounds. Therefore, it can be expected that nonstoichiometric titanium oxides

electronegative metal to the more electronegative one, and then the excess electrons are located near the more electronegative element providing preferable sites for HER [31].

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codeposited with NiW layer will influence the composite property.

Electrolytes and electrodeposition conditions. Deposition of Ni-W layer on copper plate with geometric area 2 cm^2 and carbon fibers (Fig. 1) was carried out under potentiostatic conditions (E = -1.2V and 1.0 V versus SCE) in a solution containing nickel sulphamate (Ni²⁺ 16 g/l), sodium citrate 90 g/l and Na₂WO₄.2H₂O 30 g/l. The pH of the electrolyte was 7 and the temperature 60°C [32]. 10 g/l nonstoichiometric Ti oxides (Ebonex) of Atraverda Ltd. (TiO_x particles, x = 2n-1/n, where *n* in Ti_nO_{2n-1} is ∞ , 10, 9, 8, 7, 6, 5, 4), were added and maintained in suspension by optimum stirring at 150 min⁻¹. For the electrochemical measurements in 0.5 M H₂SO₄ solution a conventional three-electrode cell (platinum electrode as a counter and SCE as a reference) was used with EG&G PAR - Versastatt, SoftCorr 352.



Fig. 1. NiW deposited layer on carbon fibers.

Characterization in acid medium. The corrosion behaviour of Ni-based composites with and without nonstoichiometric Ti oxide particles was studied by polarization curves. The deposits containing embedded TiO_x particles (Fig. 2) exhibited better corrosion behaviour [35] as the others properties retain persistent. This fact was connected with the structure of the deposits, the presence of deep cracks (Fig. 2a) as well as the amount of sorbed hydrogen.

The electrocatalytic properties of the composites were investigated by electrochemical impedance spectroscopy (EIS). The results from EIS showed that NiW + TiO_x layers on modified carbon fibre electrodes (Fig. 3a) [38] possess the highest electrocatalytic activity for HER compared with NiW + TiO_x layers on pristine carbon fiber electrodes (Fig. 3b) [38] and NiW layers on oxidized carbon fiber electrodes. This behaviour may be attributed not only to the developed electrode surface caused by the presence of nonstoichiometric titanium oxides followed by secondary crystallization of NiW layers on their surface (Fig. 2b), but also to the nature of the interaction between the composite and modified carbon fibres [38].

Codeposition of Mo

Electrolytes and electrodeposition conditions. Two types of electrolytes were used: neutral – containing nickel sulphamate (Ni²⁺ 16 g/l), C₆H₅Na₃O₇.2H₂O 90 g/l, Na₂WO₄.2H₂O 30 g/l and Na₂MoO₄.2H₂O 7.5 g/l; and a second one – NiMoW alkaline electrolyte containing Ni sulphamate (Ni²⁺ 32 g/l), C₆H₅Na₃O₇.2H₂O 72g/l, Na₂WO₄.2H₂O 24g/l, Na₂MoO₄ 6g/l, NH₄OH about 20 ml to adjust pH. The pH of the neutral electrolyte was 7 and temperature 60°C, and pH 9 and temperature 50°C for the alkaline electrolyte, respectively. The deposits from both electrolytes obtained under potentiostatic conditions (–1.0 V and –1.1 V versus SCE) were characterized with good adhesion and smoothness.



Fig. 2. Embedded TiO_x particles on NiW deposit: a) cracks; b) secondary crystallization on the TiO_x particle surface.

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Fig. 3. NiW + TiOx layers on carbon fibres and EIS measurements [38]: a) modified carbon fibre; b) pristine carbon fibre.



Fig. 4. NiMoW deposits from: a) neutral electrolyte; b) alkaline electrolyte.

Morphology and composition studies. Both deposits from neutral and alkaline electrolyte possessed globular morphology (Fig. 4). The compositions of the layers are presented in Table 1. Globular formations rich of oxygen (white balls, spectrum (Sp 1)) on the surface of deposit from neutral electrolyte were observed (Fig. 4a). Sodium-

oxide rich species (Sp 3) appeared on the surface of the deposit obtained from alkaline solution probably due to the electrolyte components (Fig. 4b). Our previous studies on the polarization curves in strongly alkaline solution for HER showed better electrocatalytic activity of NiMoW layers obtained from alkaline electrolyte than those obtained from neutral one in spite of the bigger amount of Mo in the deposit. Thus, our next studies were continued with deposits obtained from alkaline electrolyte.

Table 1. The composition of NiMoW deposits obtained from neutral and alkaline electrolyte (in weight per cent).

Neutral electrolyte							
Sp	O,	Ni,	Mo,	W,	Na,		
	%	%	%	%	%		
1	12	41	36	11	-		
2	6	47	36	11			
Alkaline electrolyte							
	0	Ni	Мо	W	Na		
3	11	58	17	4	10		
4	8	66	18	5	2		
5	5	67	22	6	-		

ELECTRODEPOSITION OF Co BASED SYSTEMS

CoMnB system

A non-conventional growth of cobalt-manganese layers deposited from electrolytes with low cobalt content was observed [39-41]. Independently on the substrate, the typical shape of the multilayered Co-Mn-B dendrite was kept, where successive growth fronts were observed, the light ones being rich of cobalt, and the dark ones - rich of oxygen, boron and manganese. Such multilayers were obtained from complex electrolyte containing 5 g/l Co²⁺, 5 g/l Mn²⁺ and 35 g/l boric acid. Cobalt electrode was used as an anode. Deposition was performed under potentiostatic conditions at -1.5 and -1.8 V (vs. SCE) on Ni-foam (RECEMAT Int. [42]), with geometric area 1 cm². The real electrochemically active area of the electrodes was calculated from curves, obtained by cyclic voltammetry in 6 M KOH electrolyte (using the Randles-Sevcik equation). The reproducible voltammograms (Fig. 5) show well defined and invariant anodic and cathodic (I_{kp}) peaks, corresponding to the formation and reduction of $Co(OH)_2$, were compared with those obtained from fresh polished and determined surface (S) of Co plate.

Growth of multi-layered dendrite formations, comprising successive layers with very high Co or O content, was observed at high deposition potentials. The obtained systems contain huge amounts of incorporated hydrogen for cobalt deposits, ca. 4200 ppm [43]. The dependence of hydrogen content remaining after thermal treatments at different temperatures suggested the presence of two types of bonded hydrogen. Based on the experimental data of the components in the different zones, and especially about the ratio between them, suggestions

were made about the nature of the BMnO layers. It was assumed that these were borate or metaborate compounds, which were adsorbed onto and then grew together with the subsequent cobalt layers. It was postulated that the presence of such layers was directly related to the absorption of considerable amounts of hydrogen in the deposits [44]. Such electrodeposited nanocomposite Co-Mn-B multi-layers on Ni-foam were investigated as catalysts for HER [30] and hydrolysis of NaBH₄ in alkaline solution [45]. It was established that the estimated value of activation energy (~55 kJ·mol⁻¹) for the tested catalysts was lower than the reported one for Co-B (~65 kJ·mol⁻¹) [46] and close to that reported for Ru-catalyst (56 kJ·mol⁻¹) [47].



Fig. 5. Reproducible voltammograms of Co electrodes in 6 M KOH.

CoNiMnB system

Sulphamate nickel (3 or 5 g/l) was added using the electrolyte described in 3.1. Thus, the concentration relation between nickel and cobalt was 3:3 g/l or 5:5 g/l.

The formation of successive light, comprising cobalt and nickel layers, and dark fronts, containing considerable amounts of oxygen and manganese, were observed by scanning electron microscopy (SEM), Fig. 6. The addition of nickel maintains typical multilayered Co-Mn-B growth. The morphology depends on concentration and electrolysis conditions.

The electrocatalytic activity of newly synthesized CoNiMnB electrodeposits for direct borohydride electroxidation was examined. Electrodes, prepared by electrodeposition from complex sulphamate electrolytes on Ni-foam at different potentials, were anodically polarized and the corresponding polarization curves were plotted. Simultaneously, the rate of generated due to borohydride hydrolysis hydrogen was monitored at each current loading. The values of overpotentials at given current densities were estimated and compared with those previously obtained with CoMnB electrodes [48].



Fig. 6. Cross-section of CoNiMnB composites obtained under potentiostatic conditions: a) E = -1.5 V vs. SCE with concentration relation between Ni and Co are 5:5 g/l; b) E = -1.8V vs. SCE with concentration relation between Ni and Co are 3:3 g/l.

 Table 2. The composition of V containing deposits (in weight per cent).

CoMnVB							
	В, %	O, %	S, %	V, %	Mn, %	Co, %	
6 7	11 18	21 39	0.2	24 31.8	15 8	29 3	
CoVB							
		0	S	V		Co	
8		38	0.3	50.7	-	11	

CoVMnB and CoVB system

Based on CoMnB electrolyte, vanadium was added as 0.5 g/l or 5 g/l VCl₃. The deposits were characterized by a huge number of cracks, worse adhesion with substrate (copper plate) than CoMnB

layers. Different morphology and vanadium content were observed (Fig. 7) depending on the electrolyte. EDX analyzes showed reach of V–O zones (Sp 7) with different morphology. Decreasing of manganese amount in the composite increased the vanadium content, too, Table 2.



Fig. 7. Vanadium codeposition with Co-based composites: a) CoVMnB; b) CoVB.

ELECTRODEPOSITION OF QUATERNARY Co AND Ni BASED SYSTEMS

Ni and its selected binary and ternary alloys with copper and cobalt over the last 10–15 years were reviewed [49]. The influence of various bath composition, pH, current density and potential conditions as well as the mechanism of electro-deposition and hydrogen evolution during electro-chemical processes were considered [20, 49]. In order to improve the electrocatalytic characteristics of this type of systems, quaternary Co-Ni based composites were synthesized.

Electrolytes and electrodeposition conditions:

CoNiMoW deposits on copper plate were produced from electrolyte containing 16 g/l Ni²⁺, 16 g/l Co²⁺ (or 3 g/l Co²⁺ for NiCoMoW deposits), C₆H₅Na₃O₇.2H₂O 72 g/l, Na₂MoO₄.2H₂O 6 g/l,

Na₂WO₄.2H₂O 24 g/l and NH₄OH to adjust pH to 10.5. The deposition was carried out under galvanostatic conditions and temperature 50°C by stirring at 200 min⁻¹. The Pt-Ti mash was used as anode and SCE – as reference electrode.

Morphology and composition studies

The influence of Co concentration and current density on the distribution of components in composites is presented in Table 3. Decreasing the initial concentration of Co about five times in the electrolyte (from 16 to 3 g/l) decreases its amount about twenty times in the deposit thus obtaining Nirich composites (NiCoMoW) as the contents of W remain nearly the same. On the other hand, the increase of current density also decreases Co amount. This fact can be explained by anomalous codeposition of iron group metals [49, 50], where the less noble metal has a stronger influence on the deposition behaviour of the partner ion. The phenomena were observed clearly on SEM images (Fig. 8). The same electrodeposition conditions but with different Co concentration in the electrolyte generated totally different morphology. The pyramidal nickel surface structures in Fig. 8a (with less amount of Co - NiCoMoW,) proceeded to small grain formations on the reach of Co composite (CoNiMoW) surface, Fig. 8b.

Table 3. Element content and thickness of the deposit *d* in Co and Ni based composites obtained under different galvanostatic conditions.

				22	2.	1000
Current	Composites	<i>d</i> ,	Ni,	Co,	W,	Mo,
density		μm	%	%	%	%
50 mA/ cm^2	NiCoMoW	9.7	69.0	11.8	3.2	16.0
70 mA/ cm^2	NiCoMoW	13.3	74.1	7.7	3.8	14.4
80 mA/cm^2	CoNiMoW	1.8	8.8	78.8	3.6	8.8
90 mA/cm^2	CoNiMoW	3.0	11.6	76.1	3.8	8.5
90 mA/cm^2	NiCoMoW	12.2	83.0	4.2	2.6	10.2
100 mA/cm^2	CoNiMoW	3.1	12.8	77.9	1.4	7.9
110 mA/cm^2	CoNiMoW	3.8	21.3	67.2	2.5	9.0
120 mA/cm^2	CoNiMoW	5.6	23.7	63.7	3.2	9.4

Polarization curves

The deposited electrodes with geometric surface 2 cm² were tested for HER in 6 M KOH. The experiments were carried out under galvanostatic steady-state conditions. Before starting the measurements, the electrode surface was polarized for 30 min by applying a relatively high (200 mA) cathodic current. Then the current was lowered to 5 mA. The reproducible polarization curves are presented in Fig. 9. The results showed that CoNiMoW and NiMoW systems possess the lowest polarization for HER compared with others, i.e. the highest electrocatalytic activity.



Fig. 8. Ni- and Co-based composites: a) NiCoMoW; b) CoNiMoW.



Fig. 9. Polarization curves of Ni- and Co-based materials.

CONCLUSION

New types of electrolytes for electrodeposition of ternary and quaternary Co- and Ni-based composites were developed. The influence of the component concentrations and electrolyte conditions on the morphology was studied. The best electrocatalytic behaviour for HER was shown by CoNiMoW and NiMoW systems. Their additional corrosion studies in different media might lead to direct application for hydrogen production and fuel cell technology.

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СИНТЕЗ И ОХАРАКТЕРИЗИРАНЕ НА ЕЛЕКТРОХИМИЧНО ОТЛОЖЕНИ НОВИ МАТЕРИАЛИ КАТАЛИЗИРАЩИ ОТДЕЛЯНЕТО НА ВОДОРОД

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(Резюме)

В настоящата работа е направен преглед на електрохимично получени собствени материали както и синтез на нови покрития от NiMoW, NiCoMnB, CoMnVB, CoVB, CoNiMoW и NiCoMoW при вариращи условия на електролизния процес. Наблюдавана е морфологията както на повърхността така и на напречни шлифове на изброените системи. Каталитичните характеристики по отношение реакцията на отделяне на водород в силно алкална среда са оценени от поляризационни зависимости. Най-добри резултати за тази реакция показват системите NiMoW и CoNiMoW. Тези материали биха могли да намерят приложение като електрокатализатори не само при производството на водород, но и в технологиите на горивните елементи.