ISSN 1729-4428

Magdalena Popczyk

The Hydrogen Evolution Reaction on Electrolytic Ni+W+Si and Ni+W+Mo+Si Composite Coatings

University of Silesia, Institute of Materials Science, Bankowa 12, 40-007 Katowice, Poland e-mail: <u>magdalena.popczyk@us.edu.pl</u>

Ni+W+Si and Ni+W+Mo+Si composite coatings were obtained by electrolytic codeposition of crystalline nickel from an electrolyte containing suspension of suitable metallic and non-metallic components (W, Mo and Si). These coatings were obtained under galvanostatic conditions, at the current density of $j_{dep.} = 0.100 \text{ A} \cdot \text{cm}^{-2}$ and at the temperature of 338 K. Chemical composition of obtained coatings was determined by EDS method. The electrochemical activity of these coatings was studied in the process of hydrogen evolution reaction (HER) from 5 M KOH solution using steady-state polarization and electrochemical impedance spectroscopy (EIS) methods. Basing on the results of EIS measurements the rate constants of the HER were determined. It was found that Ni+W+Mo+Si composite coatings are characterized by enhanced electrochemical activity towards hydrogen evolution as compared with Ni+W+Si coatings. The reason of this is connected with the presence of additional composite component (molybdenum). Thus obtained coatings may be useful in application as electrode materials for the hydrogen evolution reaction.

Key words: Nickel, Tungsten, Molybdenum, Silicon, Composite coatings, Hydrogen evolution reaction.

Стаття поступила до редакції 09.07.2007; прийнята до друку 14.09.2007.

Introduction

Many papers have dealt with ways of increasing the effectiveness of cathodes used for the hydrogen evolution reaction in alkaline solutions [1,2]. Many transition metals and metal alloys have been characterized as hydrogen electrodes. Among them, nickel and nickel-based alloys [3,4] have a high catalytic activity. However, their activity and stability are insufficient. They are usually prepared by electrolytic codeposition or dip coating. In order to improve the utilization of these materials and to enhance their electrocatalytic activity, various modifications could be applied, such as the use of alloys or composites instead of pure elements and other modifications to obtain electrodes with very developed, rough or porous electrode surface [1-4].

This study was undertaken in order to obtain the nickel coatings, containing an additional components of metallic (W) and non-metallic (Si) powders (Ni+W+Si coatings) and components of metallic (W, Mo) and non-metallic (Si) powders (Ni+W+Mo+Si coatings). Metallic (W) and non-metallic (Si) components perform a function of surface modifiers. The choice of Mo component was made based on the knowledge of properties of this metal in the hydrogen evolution reaction [3,4], which performs a function of activator in the HER. The purpose of this work was to evaluate effectiveness of these coatings as electrode materials for

hydrogen evolution in an alkaline solution.

I. Experimental

Ni+W+Si by coatings were obtained from the following electrolyte electrodeposition (concentrations in g dm⁻³): NiSO₄ \cdot 7H₂O - 84, H₃BO₃ -8, CH₃COONa - 10, C₆H₅O₇Na₃ · 2H₂O - 30, NH₄Cl - $10 + 50 \text{ g dm}^{-3}$ of tungsten powder (100 mesh, *Aldrich*) + 50 g dm⁻³ of silicon powder (325 mesh, Aldrich). were obtained Ni+W+Mo+Si coatings bv electrodeposition from the following electrolyte (concentrations in g dm⁻³): NiSO₄ \cdot 7H₂O – 84, H₃BO₃ – 8, CH₃COONa - 10, C₆H₅O₇Na₃ \cdot 2H₂O - 30, NH₄Cl -10 + 50 g dm⁻³ of tungsten powder (100 mesh, *Aldrich*) + 50 g dm⁻³ of molybdenum powder (100 mesh, Aldrich) + 50 g dm⁻³ of silicon powder (325 mesh, Aldrich). Distilled water and 'analytical grade' reagents were used for preparation of the electrolyte. The process was carried out at the temperature of 338 K, with intensive mechanical stirring (200 rpm). Coatings were electrodeposited under galvanostatic conditions, at the current density $j_{dep.} = 0.100$ A cm⁻². A platinum mesh served as an auxiliary electrode. The coatings were deposited on the steel plate (St3S) of the geometric surface area of 1 cm² and were used as working electrodes. Chemical composition of obtained coatings was determined by EDS method.

Investigations of electrolytic hydrogen evolution on the prepared coatings were conducted in a three-electrode cell, using steady-state polarization and electrochemical impedance spectroscopy (EIS) methods. These measurements were carried out in 5 M KOH solution, at temperature of 293 K, using the Autolab® electrochemical system. The auxiliary electrode was a platinum mesh and the reference electrode was of the type Hg/HgO/5 M KOH. The EIS measurements were performed approximately after 24 h of the HER at the constant current density, j = 0.320 A cm⁻². For each coating, three cycles were recorded. Each cycle consisted of 30 minutes of galvanostatic electrolysis at i = 0.320 A cm⁻² and 43 minutes of curve recording using galvanostatic polarization (45 s per point) ranging from 0.320 A cm⁻² to 0.1 μ A cm⁻². At the steady-state, the electrode potentials were corrected for iR_s drop determined by EIS. In the EIS measurements the amplitude of ac signal was equal 0.005 V. A frequency range from 10 kHz to 0.1 Hz was covered with 10 points per decade. The real and imaginary components of the complex plane plots were analysed using a modified version of a complex non-linear least-squares fitting program (CNLS) [4] from which the experimental parameters of an electrical equivalent circuit were determined. Kinetics of hydrogen evolution process was determined on basis of the dependences of both the $1/R_{ct}$ (inverse the charge-transfer resistance) and the total current density j, on overpotential ΔE . The rate constants (k_1, k_2, k_2, k_3) were evaluated using the NLS method.

II. Results and Discussion

The Ni+W+Si composite coatings contain about 39 wt% of nickel, 41 wt% of tungsten and 20 wt% of silicon. The Ni+W+Mo+Si composite coatings contain about 34 wt% of nickel, 30 wt% of tungsten, 22 wt% of molybdenum and 14 wt% of silicon. X-ray diffraction pattern showed that the deposited coatings have a composite character [5-7]. The composite is the mixture of the Ni, W and Si and mixture of the Ni, W, Mo and Si. It means that in the process of nickel electrodeposition, metallic and non-metallic powders grains have been built into a crystalline nickel matrix [5-7]. The process of powders embedding in the Ni matrix is probably based on the ability to adsorb Ni²⁺ ions on a surface of powders, which was shown earlier [8]. Such a form of nickel ions, partially devoid of its hydration envelope can be more electrochemically active than hydrated Ni^{2+} ions. A suspension of this type moves towards the cathode and by applying considerable current densities, more adsorbed Ni2+ ions than the ones hydrated in the electrolyte are submitted to the charge-transfer reaction. The composite structure is obtained on this way.

The electrochemical activity of thus obtained coatings was studied in the process of hydrogen evolution from 5 M KOH solution. The steady-state polarization curves obtained on different electrodes are presented in Figure 1 and the kinetic parameters are presented in Table 1. On the bases of steady-state of the hydrogen evolution, the rate of this process was



Fig. 1. The steady-state polarization curves obtained for HER in 5 M KOH on obtained coatings.

 Table 1

 Kinetic parameters of the HER in dependence from the

type of coating.					
Type of coating	ΔE_{100}	jo			
	[V]	$[A \text{ cm}^{-2}]$			
Ni+W+Si	-0.357	2.58 · 10-4			
Ni+W+Mo+Si	-0.266	1.03 · 10-3			

estimated depending on the type of coatings. The value of hydrogen evolution overpotential at the current density of 100 mA cm⁻² – ΔE_{100} is lower for the Ni+W+Mo+Si coatings than for the Ni+W+Si coatings. Value of exchange current density – j_0 for the Ni+W+Mo+Si coatings is higher than for the Ni+W+Si coatings. It suggests an influence of presence additional composite component (Mo) on the rate of hydrogen evolution. It was found that in the Ni+W+Mo+Si coatings the increase in the rate of hydrogen evolution was observed.

The complex impedance was analysed using a modified version CNLS fitting program [4]. Character of dependence of imaginary impedance component on electrode overpotential and real impedance component is show in the three-dimensional diagrams (3D): $-Z'' = f(Z', \Delta E)$ (Figure 2). These diagrams show influence of real impedance component and electrode overpotential on values of imaginary impedance component. The values of real and impedance components are lower for the Ni+W+Mo+Si coatings than for the Ni+W+Si coatings. This means that Ni+W+Mo+Si coatings are characterized by enhanced electrochemical activity for HER compared to Ni+W+Si coatings.

It has been found that the *ac* impedance behaviour of obtained coatings may be well described by the one-CPE electrode model. This is typical model for the HER on solid, rough or porous electrodes. Such equivalent circuit is characteristic of electrode materials composed of cylindrical pores of radius *r* and length *l*. As was shown for short and wide pores l^2/r is very small and only one semicircle on the complex plane plot was observed (model CPE1) [9].

One-CPE model consists of the solution resistance R_s in series with the parallel connection of the CPE element



Fig. 2. Dependences of $-Z^{2} = f(Z^{2}, \Delta E)$ for the Ni+W+Si coatings (a) and Ni+W+Mo+Si coatings (b).

and charge-transfer resistance R_{ct} (Figure 3) [4].

The impedance of CPE element is given as:

$$Z_{\rm CPE} = 1 / (Tj\omega)^{6}$$

where *T* is the capacitive parameter, ω is the angular frequency of *ac* voltage and ϕ is the coefficient of dispersion. As a result of approximation of experimental data in the case of CPE1 model the following parameters could be obtained: R_{ct} , *T*, ϕ , R_s .

The charge transfer resistance values, R_{ct} , are higher for the Ni+W+Si coatings than for the Ni+W+Mo+Si coatings (Figure 4). An approximation of the values of $1/R_{ct}$ in function of ΔE (in the range from -0.25 V to -0.12 V) and j in function of ΔE , was carried out by adjusting the rate constants k_1 , k_2 , k_2 , k_3 [4,10,11] and are presented in Table 2. For all obtained coatings, value of k_3 rate constant is at least of two orders smaller than values of k_1 and k_2 rate constants. Is suggesting that the main pathway for the HER at above mentioned electrode materials is Volmer - Heyrovsky, and Tafel reaction has a small contribution to this process. Values of k_1, k_{-1}, k_2 , k_{-2} and k_3 for the Ni+W+Mo+Si coatings are higher than for the Ni+W+Si coatings. This confirms that Ni+W+Mo+Si coatings characterized enhanced electrochemical activity for HER compared to Ni+W+Si coatings.

Increase in the activity of HER of Ni+W+Mo+Si coatings (higher values of j_0 , k_1 , k_{-1} , k_2 , k_{-2} , k_3 and lower values of ΔE_{100} , R_{cl}) in comparison with Ni+W+Si coatings is connected with presence of additional composite component (Mo).

Conclusions

Ni+W+Si and Ni+W+Mo+Si composite coatings



Fig. 3. Equivalent circuit scheme for the one-CPE electrode model, R_s [Ω cm²] – solution resistance, CPE – constant phase element, R_{ct} [Ω cm²] – charge-transfer resistance.

could be obtained by electrolytic codeposition of crystalline nickel with suitable metallic and non-metallic components (tungsten, molybdenum and silicon). It was found that Ni+W+Mo+Si composite coatings are characterized by enhanced electrochemical activity towards hydrogen evolution as compared with Ni+W+Si. The reason of this is connected with the presence of additional composite component (molybdenum), which performs a function of activator in the HER. Thus obtained coatings may be useful in application as



Fig. 4. Dependences of log $R_{ct} = f(\Delta E)$ for the obtained coatings.

Table 2

The rate constants of the HER in dependence from the type of coating.

Type of coating	k_{l}	<i>k</i> ₋₁	k_2	<i>k</i> ₋₂	k_3
	$[mol cm^{-2} s^{-1}]$				
Ni+W+Si	$1.68 \cdot 10^{-9}$	$1.87 \cdot 10^{-8}$	$3.96 \cdot 10^{-11}$	$3.56 \cdot 10^{-12}$	$2.47 \cdot 10^{-13}$
Ni+W+Mo+Si	$3.04 \cdot 10^{-9}$	$5.65 \cdot 10^{-8}$	$1.35 \cdot 10^{-10}$	$7.27 \cdot 10^{-12}$	$9.11 \cdot 10^{-13}$

electrode materials for the hydrogen evolution reaction in *T08C 009 28* comparison with Ni+W+Si coatings.

This research was partially financed with Project 3

- [1] M. Popczyk, A. Budniok, A. Lasia. Electrochemical properties of Ni-P electrode materials modified with nickel oxide and metallic cobalt powders *Int. J. Hydrogen Energy* **30**, pp. 265-271 (2005).
- [2] M. Popczyk, A. Serek, A. Budniok. Production and properties of composite layers based on an Ni-P amorphous matrix *Nanotechnology* 14, pp. 341-346 (2003).
- [3] J. Niedbała, M. Popczyk, J. Panek, A. Budniok, E. Łagiewka. Influence of thermal treatment on process of hydrogen evolution on electrolytic nickel-molybdenum layers *Inżynieria Materiałowa* 1(144), pp. 31-34 (2005).
- [4] R.K. Shervedani, A. Lasia. Study of the hydrogen evolution reaction on Ni-Mo-P electrodes in alkaline solutions J. Electrochem. Soc. 145(7), pp. 2219-2225 (1998).
- [5] M. Popczyk, J. Kubisztal, A. Budniok. Structure and electrochemical characterization of electrolytic Ni+Mo+Si composite coatings in an alkaline solution *Electrochim. Acta* **51**, pp. 6140-6144 (2006).
- [6] M. Popczyk, A. Budniok, E. Łągiewka. Structure and corrosion resistance of nickel coatings containing tungsten and silicon powders *Mater. Charact.* 58, pp. 371-375 (2007).
- [7] M. Popczyk, J. Kubisztal, A. Budniok. Electrodeposition and thermal treatment of nickel coatings containing molybdenum and silicon *Mater. Sci. Forum* 514-516, pp. 1182-1185 (2006).
- [8] A. Serek, A. Budniok. Electrodeposition and thermal treatment of nickel layers containing titanium *J. Alloys Compd.* **352**, pp. 290-295 (2003).
- [9] A. Lasia. Nature of the two semi-circles observed on the complex plane plots on porous electrodes in the presence of a concentration gradient *J. Electroanal. Chem.* **500**, pp. 30-35 (2001).
- [10] J.M. Jakšić, M.V. Vojnović, N.V. Krstajić. Kinetic analysis of hydrogen evolution at Ni-Mo alloy electrodes *Electrochim. Acta* 45, pp. 4151-4158 (2000).
- [11] S. Rodrigues, N. Minichandraiah, A.K. Shukla. Kinetics of hydrogen evolution reaction on Zr_{0.5}Ti_{0.5}V_{0.6}Cr_{0.2}Ni_{1.2} alloy in KOH electrolyte *Bull. Mater. Sci.* 23(5), pp. 383-391 (2000).

Магдалена Попчик

Реакція водневої еволюції в електролітичних композитних сумішах Ni+W+Si та Ni+W+Mo+Si

Університет Сілесія, Інститут матеріалознавства, 12, вул.. Банкова, 40-007 м. Катовіце, Польща, E-mail: <u>magdalena.popczyk@us.edu.pl</u>

Композитні суміші Ni+W+Si і Ni+W+Mo+Si були одержані електролітичним осадженням кристалічного нікелю з електроліту, що містить домішки відповідних металевих і нерудних компонентів (W, Mo i Ci). Ці суміші були одержані в гальваностатичним осадженням, при густині струму $j_{dep.} = 0,100$ cm⁻² і при температурі 338 К. Хімічний склад одержаних сумішей був визначений методом EDS. Електрохімічна діяльність цих сумішей вивчалася в процесі реакції водневої еволюції. Знайшли, що суміш Ni+W+Mo+Si характеризуються збільшеною електрохімічною діяльністю у напрямку до водневої еволюції порівнянно з сумішшю Ni+W+Si. Причина цього пов'язана з присутністю додаткового складового компоненту (молібден).

Ключові слова: нікель, вольфрам, молібден, кремній, композитні суміші, реакція водневої еволюції.