Synthesis and functional properties of mixed titanium and cobalt oxides

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Peculiarities of plasma-electrolytic oxidation of the aluminum alloys in diphosphate electrolytes are discussed. It is shown that PEO parameters depend on the composition and concentration of the components of the working solutions. The mechanism of cobalt oxides incorporation into the composition of surface layers is proposed. It has been established that the oxidation of titanium in citrate-diphosphate electrolytes promotes the formation of mixed oxide layers TiO₂-CoOₓ. The obtained mixed oxide systems have a developed microporous globular-toroidal surface and are characterized by a complex of enhanced functional properties — corrosion and abrasion resistance, catalytic activity in the carbon (II) oxide conversion reaction.

Keywords: titanium alloy, plasma electrolytic oxidation, titania, cobalt oxides, mixed oxides, corrosion resistance, catalytic activity, abrasion resistance.

Рассмотрены особенности плаズменно-электролитического оксидирования сплавов титана в щелочных билигандных электролитах. Показано, что параметры ПО зависят от состава и концентрации компонентов рабочих растворов. Предложен механизм инкорпорации оксидов кобальта в состав поверхностных слоев. Установлено, что оксидирование титана в цитратно-дифосфатных электролитах способствует формированию смешанных оксидных слоев TiO₂-CoOₓ. Полученные смешанные оксидные системы имеют развитую микропористую глобуллярно-торoidalную поверхность и характеризуются комплексом повышенных функциональных свойств — коррозионной стойкостью, стойкостью к абразивному износу, катализитической активностью в реакции конверсии оксида углерода (II).

Синтез та функціональні властивості змішаних оксидів титану та кобальту. М.В.Вед'ї, М.Д.Сахненко, Г.В.Каракурчі. М.В.Майба, О.В.Галак.

Розглянуто особливості плаズмено-електролітичного оксидування сплавів титану в лужних білігандних електролітах. Показано, що параметри ПЕО залежать від складу і концентрації компонентів робочих розчинів. Запропоновано механізм інкорпорації оксидів кобальту до складу поверхневих шарів. Встановлено, що оксидування титану у цитратно-дифосфатних електролітах сприяє формуванню змішаних оксидних шарів TiO₂-CoOₓ. Отримані змішані оксидні системи мають розвинену мікропористу глобуллярно-торіобранину поверхню і характеризуються комплексом підвищених функціональних властивостей — корозійною стійкістю, стійкістю до абразивного зношування, каталітичною активністю в реакції конверсії оксиду карбону (II).

1. Introduction

Many unique properties of titanium and titanium-based alloys specify the demand for it and its broad application by the medicine, utility companies and other branches of the industrial sector. In comparison with other structural materials titanium has two main advantages, in particular high specific strength and corrosion resistance even in
the harsh environment and when in contact with many aggressive chemical reagents [1]. Other properties of titanium, for example non-magneticity, high melting temperature, low coefficient of thermal expansion, etc. are of great importance for other application fields [2–4]. Its biological inertness in the human body, high mechanical and anti-corrosion properties specify its wide application for the medicine. Titanium and its alloys are characterized by a high resistance to the fatigue wear at alternate loads, which is rather important for the manufacture of intraosseal locks, internal and external prosthetic devices [5–7].

At the present time, many techniques and methods are available that are used for the formation of functional coatings at titanium alloys that enable the expansion of the spectrum of physical and mechanical properties of the materials and the field of their application [8, 9]. Electrochemical technique for the synthesis of oxide layers is one of the most practically feasible and economically efficient methods because it enables the production of coatings uniform in thickness and composition with high adhesive properties [10–13].

As of today, an ample amount of methods are available that are used for the formation of functional coatings doped with different components, in particular transition metal oxides. However most of them are characterized by high labor-intensive-ness and multistage process [14, 15]. Providing a uniform sediment distribution and sufficient adhesion presents a considerable problem. The method of plasma-electrolytic oxidation enables the formation of oxide matrix-substrate and the filling of it with the transition metal oxide within one stage, and as a result, a high dopant content is obtained, which is uniform in thickness and composition, and a high adhesive strength is also achieved [16].

Titanium like other valve metals is subjected to the oxidation to obtain coatings of a different purpose, for example these can be used as catalytically active layers, corrosion resistant and antifriction coatings, etc. The guided synthesis of the films of a prescribed composition and structure is required to create efficient protective coatings and new materials. The coatings formed in the microplasma mode are characterized by the biological inertness, X-ray transparent, corrosion resistance and the developed globular structure. It is known that metal oxides are used as catalysts for many technological processes in gas and liquid media, and as electrode materials for electro-synthesis and chemical current sources, etc. [17]. The oxides of non-stoichiometric composition are of special interest, because the rate of electrochemical and chemical processes is increased with an increase in a degree of deviation from the stoichiometry [18]. In this connection it is worth to note that the method of plasma-electrolytic oxidation (PEO) allows us to improve physical and chemical properties of the items, create universal polyfunctional tissue-compatible composite materials, imparting appropriate properties to them, in particular anticorrosion, tribological and bioinert properties and get the contact surface of implanted structures of a wanted structure and thickness [19–21].

Developing new approaches to the formation of surface structures on the metals and alloys that contain the specified ratio of the compounds of bivalent, trivalent and polyvalent metals, including precious metals is of great interest for the creation of new functional materials used for the catalysis, for example for the processes that require high heat conductivity [22]. In this connection, the doping of titanium oxides with cobalt is considered to be the most promising approach to the synthesis of catalytic and biocompatible materials [23].

Peculiarities of the PEO synthesis of mixed oxide coatings TiO_xCoO_y and their functional properties are considered in this article.

2. Experimental

The composition of studying titanium VT1-0 alloy was, wt.%: Ti — 99.2–99.7, Fe < 0.25, C < 0.07, Si < 0.1, N < 0.04, O < 0.2, and impurities — 0.3. The pre-treatment of the samples included mechanical cleaning from the process contaminants, degreasing in a 0.2–0.3 M NaOH, etching in a mixture of a 0.1–0.3 M HF and 0.3–0.9 M HNO_3, and rinsing with distilled water.

Coatings were formed in an electrolyte composition, mol dm−3: 0.3 K_3P_2O_7, 0.1 Na_2C_2O_4H_2O_7, 0.1 CoSO_4.

The PEO of VT1-0 alloy was conducted using a B5-50 stabilized power supply that maintained voltage up to 300 V. PEO was implemented in an electrolytic cell under conditions of forced cooling of the electrolyte to a temperature of 25–30°C and stirring the electrolyte. A current density varied in a range of 1–5 A dm⁻², the processing time was 30 to 60 min.
The chemical composition of the coatings was determined by X-ray photoelectron spectroscopy on an INCA Energy 350 energy dispersive spectrometer; X-ray radiation was excited via exposing the samples to a 15 keV electron beam. In addition, X-ray fluorescence analysis was conducted using a SPRUT portable spectrometer with a relative standard deviation of 10^{-3}–10^{-2}; the error in determining the component content was ±1 wt. %.

The surface morphology of the coatings was studied using a Zeiss EVO 40XVP scanning electron microscope (SEM). Images were recorded by the registration of secondary electrons (BSEs) via scanning with an electron beam; this mode made it possible to study the topography with a high resolution and contrast ratio.

The structure of the deposits was examined by X-ray diffraction analysis using a diffractometer (DRON-2.0) in the emission of iron anode and CuKα radiation.

The corrosion behavior of titanium samples with oxide coatings in aggressive test medium 0.1 M NaCl was studied by impedance spectroscopy technique [24]. Measurements were performed using automatic alternating current bridge R-5083 over the serial equivalent circuit in frequency range 20–1\times10^5 Hz. Auxiliary electrodes — coplanar plates of corrosion-resistant steel X18H10T. Potentiometric measurements in media 2M NaOH and Ringer’s (composition, mol dm^{-3}: NaCl — 0.15, KCl — 0.004, CaCl_{2} — 0.004) solutions were used to verify the results.

The abrasion rate for the materials \Delta l/\Delta m was determined by the Calotte method on the calotester [25].

The catalytic properties of the oxide systems were studied in the model reaction oxidation of carbon (II) oxide to carbon (IV) oxide as in [26].

3. Results and discussion

Chromatograms of the voltage of mixed oxide coatings (Fig. 1a) have a classic shape with three typical sections. In the pre-spark region (section I) U(t) dependences obtained in diphosphate electrolyte (Fig. 1a, curve 1) are actually linear due to the formation in first 2–3 min of the barrier titanium oxide with the current efficiency close to 100 %. In the electrolytes that contain Co(II), the voltage at section I is higher (Fig. 1a, curve 2), consequently Co(II) oxidation and formation of mixed TiO_2-Co_3O_4 oxides occur already in the pre-spark region. It should be noted that the cobalt oxidation (II) can progress by the following routes:

\begin{align}
3\text{Co(OH)}_2 + 2\text{OH}^- &= \text{Co}_3\text{O}_4 + 4\text{H}_2\text{O} + 2\text{e}, \\
2\text{Co(OH)}_2 + 2\text{OH}^- &= \text{Co}_2\text{O}_3 + 3\text{H}_2\text{O} + 2\text{e}, \\
\text{Co(OH)}_2 + 2\text{OH}^- &= \text{CoO}_2 + 2\text{H}_2\text{O} + 2\text{e}, \\
2\text{Co}_3\text{O}_4 + \text{H}_2\text{O} &= 3\text{Co}_2\text{O}_3 + 2\text{H}^+ + 2\text{e}, \\
\text{Co}_2\text{O}_3 + \text{H}_2\text{O} &= 2\text{CoO}_2 + 2\text{H}^+ + 2\text{e}.
\end{align}

The reactions (1–5) result in the formation of insufficiently stable mixed cobalt oxide (II, III) that crystallizes in the spinel lattice with the specific resistance of (1.05–1.6)\times10^4 \Omega \cdot \text{cm}. The CoO oxide with higher specific electric resistance 10^6–10^{10} \Omega \cdot \text{cm} is formed in the discharge channel at temperatures above 1230 K by the reaction:
Table 1. Ti\textsuperscript{III}O\textsubscript{3}Co\textsubscript{y} system corrosion indices in the 0.1 M NaCl solution

<table>
<thead>
<tr>
<th>Cobalt content in the coating, at.%</th>
<th>Corrosion depth index ( k_i \times 10^4 ) \text{mm/year}(^{-1} )</th>
<th>Corrosion potential ( E_{\text{corr}} ) V</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>2.33</td>
<td>0.05</td>
</tr>
<tr>
<td>3</td>
<td>1.55</td>
<td>0.061</td>
</tr>
<tr>
<td>5</td>
<td>1.16</td>
<td>0.094</td>
</tr>
<tr>
<td>10</td>
<td>0.99</td>
<td>0.159</td>
</tr>
</tbody>
</table>

\[ 2\text{Co}_3\text{O}_4 \rightarrow 6\text{CoO} + \text{O}_2, \quad \Delta H^0_{298} = 322.8 \text{ kJ} \quad (6) \]

With the origination of sparking (section II) an increase in the voltage is considerably retarded (Fig. 1b) due to the breakdown of the barrier film TiO\textsubscript{2}. The titanium oxide formation rate is reduced and cobalt oxides Co\textsubscript{3}O\textsubscript{4} are subjected to the thermal decomposition:

\[ \text{CoO}_2 \rightarrow \text{Co}_2\text{O}_3 \rightarrow \text{Co}_3\text{O}_4. \quad (7) \]

The transition to the region of microarcs (section III) is characterized by an insignificant change in the voltage and relative process stability (Fig. 1b), and PEO voltages are in the range of 120 to 160 V. The sparking voltage shows the slope opposition to the cobalt concentration in the oxide coating (Fig. 1c), which is due to the increase in the content of cobalt in the oxide coating.

The data of microscopic studies of the morphology of oxide coatings are indicative of the formation of tubular surface TiO\textsubscript{2}, (Fig. 2a, b) and torus-like surface TiO\textsubscript{2}Co\textsubscript{y} with a rather lower porosity (Fig. 2c, d). Such a structure may provide the good substrate for the body cells deposition. In the case of artificial hip joint, the crucial parameter is a good sliding ability, which may be provided by the surface of a low roughness.

The distribution of main elements across the thickness of TiO\textsubscript{2}Co\textsubscript{y} coating (Fig. 3) proves the fact of inclusion of alloyed metals into the titanium oxide matrix, which can be achieved particularly in the PEO mode. A prevalence of titanium at the Ti|oxide boundary and an increase in the concentration of phosphorus inclusions on the coating surface are quite natural.

A change in the behavior of diffraction patterns and also in the intensity of titanium oxide peaks and the origination of cobalt oxide peaks (Fig. 4) are indicative of the formation of mixed oxides TiO\textsubscript{2}Co\textsubscript{y} [27].

Impedance plots of mixed metal oxide systems with different cobalt content have
the same shape. At high frequencies experimental points belong to semicircumference elements and at low frequencies the linear section appears with the slope angle of about 45° that corresponds to the Warburg impedance (Fig. 5a). Such character of diagrams enables the description of metal oxide systems using the equivalent circuit (Fig. 5b) that contains the elements with the specific physical meaning: $R_E$ is the electrolyte resistance, $C_d$ corresponds to the double electric layer capacitance at low frequencies and parametric resistance elements $R_f$ and $Z$ reflect the final rate of electrochemical reaction and that of the diffusion and their interdependence. The resistance $R_f$ is in inverse proportion to the current corrosion index, therefore it can be used for the determination of the corrosion resistance of materials [28].

![Graph](image1)

**Fig. 3.** Thickness distribution of the main components in the PEO-coating TiO$_x$CoO$_y$.

**Fig. 4.** Diffraction pattern of PEO coatings consisting of mixed oxides TiO$_x$CoO$_y$.

The corrosion depth index $k_d$ (Table 1) defined using $R_f$ showed that the corrosion resistance of TiO$_x$CoO$_y$ systems is increased with an increase in the content of cobalt in the coating.

The corrosion test data for TiO$_x$CoO$_y$ coatings in 2 M NaOH and Ringer’s solutions (Table 2) are also indicative of high corrosion resistance.

In the case of TiO$_x$CoO$_y$ system, the highest anodic value of the corrosion potential may be associated with the close packed doughnut like grains filled with the small grains. The lowest corrosion resistance of coating TiO$_x$ may be associated with the cracking of the formed layer, which promotes the penetration of electrolyte to the substrate.

The most significant is the resistance of mixed oxide systems TiO$_x$CoO$_y$ to the abrasive wear. The abrasive wear rate $\Delta I/\Delta t$ of the titania coating of composition, wt. %: O

![Graph](image2)

**Fig. 5.** Impedance plots (a) and the equivalent circuit (b) of oxide coatings TiO$_x$CoO$_y$ with the cobalt content of at.%: 1 — 10; 2 — 5; 3 — 3; 4 — 0.

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Table 2. Corrosion potentials of the specimens with the oxide coating TiO₂-CoO₃

<table>
<thead>
<tr>
<th>Electrolyte composition for</th>
<th>Micrograph of the surface</th>
<th>Corrosion potential</th>
<th>Ecorr, V</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEO, mol dm⁻³</td>
<td>x1000</td>
<td>2M NaOH</td>
<td>Ringer's solution</td>
</tr>
<tr>
<td>K₂PO₄ - 0.3</td>
<td></td>
<td></td>
<td>0.4</td>
</tr>
<tr>
<td>NaC₆H₅O₇-0.1</td>
<td></td>
<td></td>
<td>0.6</td>
</tr>
<tr>
<td>CoSO₄-0.1</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- 48.22; Ti — 28.73; P — 16.25; K — 6.81, and 30 μm thickness is 1.05 μm⁻¹. Even at a coating thickness of 26 μm a decrease in the diamantine abradion rate for cobalt containing oxide system (O — 58.22; Ti — 29.02; Co — 21.11; P — 7.95; K — 2.7) up to 0.75 μm⁻¹ is observed. Such behavior may be explained due to the following reasons; in particular uniform distribution of cobalt over the coating thickness (see Fig. 3), globular-tubular structure and the spherical shape of grain agglomerates (Table 2).

Test data of the catalytic activity of coatings consisting of cobalt-based mixed oxides TiO₂-CoO₃ with α(Co) = 5–7 wt. % that were tested using the model oxidation reaction of carbon oxide (II) show that the ignition temperature Tᵢ that corresponds to the start of the efficient work of catalyst is in the range of 250 to 270°C and it is somewhat higher than that of platinum contacts (200°C). At 420°C TiO₂-CoO₃ coatings provide the CO conversion level of 68 % (for Pt contacts — 100 %), and the full conversion is reached at higher temperatures. Such activity of oxide systems containing cobalt is quite acceptable for the creation on their basis of catalytic converters of toxicants of various natures.

4. Conclusions

Plasma electrolytic treatment of titanium alloys in citrate-diphosphate electrolyte at a voltage of 120 to 160 V enables the formation of mixed metal oxide systems with the cobalt content of 5 to 10 at. %. It is shown that the availability of inverse relation of the sparking voltage as a function of the dopant concentration in the electrolyte is conditioned by an increase in the electric conductivity of mixed oxides that gain thickness due to a higher conductivity of oxide dopant compounds.

TiO₂-CoO₃ coatings have a microporous globular torus-like surface without cracks that provides both substantial corrosion and high abrasive wear resistance.

Mixed oxide coatings are characterized by the developed surface and high catalytic activity in the carbon (II) oxide conversion reaction. This confirms the efficiency of materials as catalysts for the gaseous waste neutralization and the processes of hydrocarbon combustion.

References