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## COMPARATIVE ANALYSIS OF ELECTROCHROMIC PROPERTIES OF CuWO<sub>4</sub>•WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> AND WO<sub>3</sub> THIN FILMS

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**Abstract:** A comparative analysis of electrochromic properties of composites  $CuWO_4 \bullet WO_3$ ,  $Bi_2WO_6 \bullet WO_3$  and  $WO_3$  films obtained by electrochemical and chemical methods was carried out. The study into the kinetics of light transmission and spectral characteristics of electrochromic coloration revealed some differences in electrochromic processes. It found that in the  $WO_3$ ,  $Bi_2WO_6 \bullet WO_3$ ,  $CuWO_4 \bullet WO_3$  series, lithium intercalation in the film is slowed down, which is due to diffusion limitations in the process of coloring of the Bi and Cu oxides. Spectral characteristics of light transmission  $Bi_2WO_6 \bullet WO_3$  and  $CuWO_4 \bullet WO_3$  also differ from  $WO_3$  in that the contribution to light absorption is also made by Bi and Cu oxides, which are partially reduced by lithium in the process of their coloring. It is shown that the metal tungstates can be effective electrochromic materials with an additional absorption band in the visible region.

**Keywords:** electrochromism, metal tungstates, electrochromic composites.

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#### Introduction

There is not enough published studies about electrochromic properties of metal tungstates what delays their use in real devices as smart windows, multicolor bistable displays, solar heat regulation, optical telecommunications and applications in aerospace and military camouflage [1-5]. In this article, thin films of CuWO<sub>4</sub>•WO<sub>3</sub> Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> were selected as objects of research into the electrochromic properties of metal tungstates. The tungstates of these metals can have similar spectral characteristics of electrochromic coloration to tungsten oxide with some differences typical for these materials. For the synthesis of films, preference was given to electrodeposition methods with interference control of film thickness. According to the analysis, literature the method electrochemical deposition provides films with a high degree of hydration [6, 7] and allows controlling the thickness [8] and surface morphology [9] of films, while the choice of the optimal thickness in the range of 150 - 200 nm [10, 11] allows, in turn, expecting the most optimal parameters of electrochromic efficiency and coloration rate.

#### **Experimental part**

 $\text{CuWO}_4\text{-}\text{WO}_3$  films were obtained in two stages by  $\text{Cu}_2\text{O}$  electrochemical deposition at the cathode current 1 mA/cm² from solution of the composition (CuSO<sub>4</sub>- 0.05 mol/l, citric acid and 2 mol/l KOH, pH = 10) and followed by anodized Cu<sub>2</sub>O in a solution of 1 mol/l K<sub>2</sub>WO<sub>4</sub> at a voltage of 3.5 V for 0.5 hours. The

peroxide electrolyte  $Na_2WO_4$ -0.1 mol/l,  $H_2O_2$ -0.2 mol/l and  $H_2SO_4$  (pH = 1.1) was used to obtain  $WO_3$  film. The deposition was carried out in the galvanostatic mode with a cathodic current of 1.5 mA/cm<sup>2</sup>. Bismuth tungstate was taken off by electrochemical precipitation from an electrolyte containing  $Bi_2O_3$ = 45 g/l,

 $Na_2WO_4 = 100$  g/l, 35%  $H_2O_2$  - 50 ml/l, adjusted to pH = 1 with nitric acid (55 ml) at a cathodic current of 1 mA/cm². Bismuth tungstate was also obtained by ion layering using solutions of  $Bi_2O_3 = 45$  g/l, pH = 1 (nitric acid 55 ml) and  $Na_2WO_4 = 100$  g/l. All used chemical reagents were analytical grade purchased from Sigma-Aldrich. The XRD study of films was performed using a DRON 4 diffractometer. The optical properties of the films were studied using a Perkin Elmer UV – Vis Lambda 35 spectrophotometer.

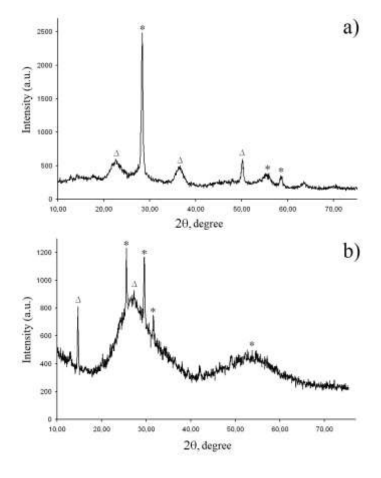
The electrochemical measurements of deposited films were performed using PGSTAT Elins P-8S Potentiostat. Platinum was used as a counter electrode and  $SnO_2$  / glass substrate as working electrode. The EC study was carried

out in 1 M LiClO<sub>4</sub> in propylene carbonate solutions. The change in the light transmittance (electrochromic color) of the films measured using a universal setup based on a single-beam diffraction spectrophotometer of the C-302 type, which provided measurements in the wavelength range  $\lambda$  from 300 to 1300 nm. The monochromator was controlled using a complex based on a personal computer. The galvanostatic current change was provided with the help of the G6-26 signal generator, which supplies a current control pulse to potentiostat and electrochromic cell. At the same time, the dynamics of changes in the intensity of the transmitted light through the film were recorded.

#### **Results and discussion**

The mechanism of electrochemical formation of Bi<sub>2</sub>WO<sub>6</sub> films is similar to the processes of formation of WO3 as a result of of peroxide-complex electroreduction compounds based on tungstate ions. The interaction of Na<sub>2</sub>WO4 and H<sub>2</sub>O<sub>2</sub> forms a  $[(O_2)_2(O)W-O$ complex peroxotungstate  $W(O)(O_2)_2^{2-1}$  [12]. The work [13] shows that the process of electrodeposition of WO<sub>3</sub> from acidic solutions containing this complex is carried out in two stages: 1 - electrochemical breaking of the O-O bond in the molecule of the peroxotungstate complex and 2 – the chemical stage of polymerization to tungstic acid that forms tungstate ions (H<sub>2</sub>WO<sub>4</sub>=WO<sub>4</sub><sup>2-</sup>+2H<sup>+</sup>) and is present in the near-cathode space. If ions (Bi<sup>3+</sup>) are added to the deposition solution, along with the formation of H<sub>2</sub>WO<sub>4</sub> the Bi<sub>2</sub>WO<sub>6</sub> will also be co-precipitated. Studies of the structure of the obtained films proved this assumption. From the interpretation of X-rays, fig. 1, it is established that along with Bi<sub>2</sub>WO<sub>6</sub>, hydrated phases of WO<sub>3</sub> are also observed. Fig. 1 presents the results of X-ray phase analysis of bismuth tungstate obtained by ion layering (a) electrochemical method (b). and X-ray diffraction analysis established that the composition of the materials is mixed and includes, in addition to bismuth orthorhombic tungstate, tungsten trioxide with a hexagonal

structure. The comparison of X-ray phase analysis for chemically and electrochemically obtained Bi<sub>2</sub>WO<sub>6</sub> showed that the films obtained by electrochemical deposition have an structure, with amorphous interspersed crystallites of orthorhombic Bi<sub>2</sub>WO<sub>6</sub> hexagonal WO<sub>3</sub> (Fig. 1 (b)) [14, 15], while the films obtained by ion layering have a polycrystalline structure with broadened peaks (Fig. 1 (a), which indicates the fine-grained nature of the obtained crystallites. This is explained as being due to the fact that during ion layering, crystal points are created for further crystal growth, and during electrochemical deposition, a process similar to polymerization takes place following which an amorphous mixture of mixed phase composition Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> is formed. To obtain thin CuWO<sub>4</sub>•WO<sub>3</sub> films with optimal electrochromic parameters [10, 11], voltammetric studies of Cu<sub>2</sub>O electrodeposition as a precursor for the formation of CuWO<sub>4</sub>•WO<sub>3</sub> were performed. The choice of Cu<sub>2</sub>O current electrodeposition is based on voltammetric studies. Fig. 2 shows voltammetry of the Cu<sub>2</sub>O precipitation process from citrate solution based on CuSO<sub>4</sub> and alkali. From the analysis of the curve of Fig. 2 it was found that within the potentials of the reduction wave from Cu<sup>2+</sup> to Cu<sup>+</sup> [16] the current is 1 mA/cm<sup>2</sup>. As a result, Cu<sub>2</sub>O films are formed.



**Fig.1.** a - XRD pattern of chemically obtained bismuth tungstate, where \*- orthorhombic  $Bi_2WO_6$ ,  $\Delta$  - hexagonal  $WO_3$ ; b - X-ray phase analysis of electrochemically obtained bismuth tungstate.

The resulting  $Cu_2O$  films were anodized in 1 mol / 1  $K_2WO_4$  solution at 3.5 V for 30 min. The anodization resulted in the dissolution of

Cu<sub>2</sub>O and the formation of CuWO<sub>4</sub> sediment. The process of anodizing Cu<sub>2</sub>O can be described by reactions [17] as follows:

$$Cu_2O + H_2O = 2Cu^{2+} + 2OH^{-} + 2e^{-}$$
 (1)

$$WO_4^{2-} + Cu^{2+} = CuWO_4$$
 (2)

Upon completion of these reactions, thin films of  $CuWO_4$  copper tungstate with  $WO_3$  impurities were obtained. In parallel with the  $CuWO_4$  formation there is the reaction of water

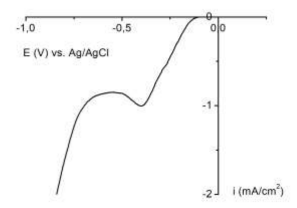
decomposition at the anode with the release of oxygen and  $H^+$  protons that interact with  $WO_4^{2-}$  to form hydrated forms of  $WO_3$ :

$$2H_2O = 4H^+ + O_2 + 4e^-$$
 (3)

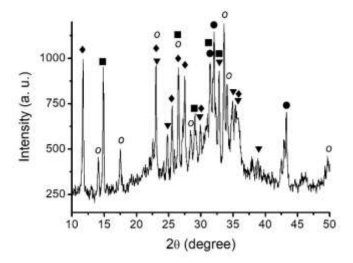
$$WO_4^{2-} + 2H^+ = WO_3 \cdot H_2O$$
 (4)

X-ray phase analysis CuWO<sub>4</sub>•WO<sub>3</sub> on the Fig. 3 showed the presence of monoclinic structure CuWO<sub>4</sub> • 2H<sub>2</sub>O (standard card (PDF 33-0503)) [18,19] and undeciphered tungstate structure, possibly, phase CuWO<sub>4</sub> (standard card (JCPDS No 88-0269)) [20] with impurities of

hydrated forms  $WO_3$  monoclinic  $WO_3 \cdot 2H_2O$  (JCPDS Card No.18-1420) [21], orthorhombic  $WO_3 \cdot 0.33H_2O$  (JCPDS Card No. 35-0270) [22] and orthorhombic structure  $WO_3$ , (JCPDS card 20-1324) [23].



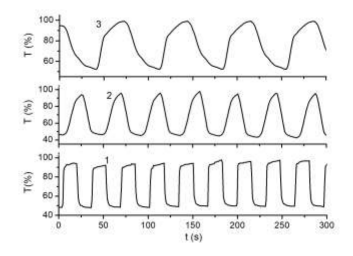
**Fig. 2.** Voltammetry of the process of obtaining  $Cu_2O$  from a solution ( $CuSO_4$ - 0.05 mol / l, citric acid 2 mol / l, KOH -to pH = 10). Scan rate 5 mV/s.



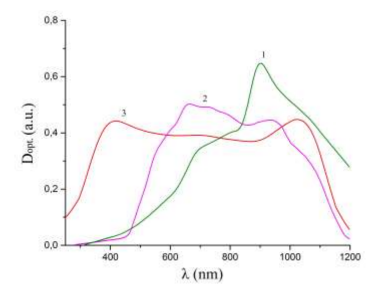
**Fig. 3**. XRD pattern of CuWO<sub>4</sub>•WO<sub>3</sub> sample where ■- hydrated form of monoclinic CuWO<sub>4</sub>·2H<sub>2</sub>O, ● - undeciphered tungstate structure [20], ▼- monoclinic WO<sub>3</sub>•2H<sub>2</sub>O, o- orthorhombic WO<sub>3</sub>•0.33H<sub>2</sub>O, ■- orthorhombic structure tungsten trioxide

To assess the stability of the obtained bismuth and copper tungstate films as an electrochromic material, they were cycled in the galvanostatic mode with a current from +2.5 to -2.5 mA/cm<sup>2</sup> in 1 M LiClO<sub>4</sub> in propylene carbonate solutions while measuring light transmission (Fig. 4). The comparison of the cycling rate of bismuth and copper tungstates with tungsten trioxide showed different rates of activity for the samples. Also, the comparison of the kinetics of color change processes of electrochromic films of copper and bismuth tungstates with tungsten trioxide showed that in the WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub>, CuWO<sub>4</sub>•WO<sub>3</sub> series, lithium intercalation in the films slows down. As can be seen from Fig. 4, the change in the light transmittance over the same period of time in three samples slows down in the series WO<sub>3</sub> (curve 1),  $Bi_2WO_6 \cdot WO_3$  (curve 2) and CuWO<sub>4</sub>•WO<sub>3</sub> (curve 3), which is due to the mobility of ions in these structures [24]. In the series WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> and CuWO<sub>4</sub>•WO<sub>3</sub>, the highest mobility of charge carriers is observed in amorphous WO<sub>3</sub> films of 20 cm<sup>2</sup> /V.s [24], the lowest in  $CuWO_4$  0.006 cm<sup>2</sup> /V.s [25]. Since the effect of an electric field accelerates electrochromic processes only in those materials in which there are no significant obstacles to the intercalation of ions and the injection of electrons into the films, the materials whose composition includes oxides with the lowest mobility of charge carriers will have the lowest lithium diffusion rate. At the same time, in the films of Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> and CuWO<sub>4</sub>•WO<sub>3</sub> tungstates, a partial reverse reduction of oxide compounds of bismuth and copper to lower oxides is observed in

comparison to tungsten trioxide. The results obtained show that differences in spectral characteristics are observed in composite films, which is expressed by the shift of the absorption band to the region of shorter wavelengths.



**Fig. 4.** Cyclic dependences of light transmittance for WO<sub>3</sub> films (1), Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> (2) CuWO<sub>4</sub>•WO<sub>3</sub> (3) in the galvanostatic mode with a current of 2.5 and -2.5 (mA/cm2) ( $\lambda = 1000$  nm)



**Fig. 5.** Optical spectra of electrochromic coloration for WO<sub>3</sub> (1) Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> (2) CuWO<sub>4</sub>•WO<sub>3</sub> (3) at j=-2.5 mA/cm<sup>2</sup> in 1 M LiClO<sub>4</sub> in propylene carbonate solution.

Fig. 5 shows the spectral characteristics of light transmittance of colored films of copper bismuth tungstates and tungsten trioxide. From Fig. 5 curve 1 it follows that for tungsten trioxide the absorption maximum is within  $\lambda$ =1000 nm. For bismuth tungstate obtained by electrodeposition, the absorption maximum occurs at 650 nm. And for CuWO<sub>4</sub> film with lithium intercalation, an even greater shift of the

absorption band to  $\lambda$ =400 nm is observed. The comparison of the spectral characteristics of CuWO<sub>4</sub>•WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> and WO<sub>3</sub> showed that other oxide components besides WO<sub>3</sub> contribute to the color of the film. The analysis of literature data found that the transition from Cu<sup>2+</sup> to Cu<sup>+</sup> in copper oxide compounds is accompanied by the appearance of an absorption band at 400 nm [26] as in Fig. 4,

curve 3. And the partial reversible electrochemical reduction of Bi<sub>2</sub>O<sub>3</sub> during Li<sup>+</sup> intercalation leads to the decrease in the transmission (increase in absorption) of light in the 550-650 nm range [27], which is also observed in our case in fig. 5, curve 2. Thus, in addition to the absorption band of tungsten

trioxide, which falls mainly on the IR region (Fig. 5, curve 1), additional absorption bands appear in the visible part of the spectrum. This fact reveals that the use of metal tungstates is promising, provided that the intercalation of lithium into oxide components is sufficiently large.

#### **Conclusions**

By using of the combined chemical and electrochemical methods WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> and CuWO<sub>4</sub>•WO<sub>3</sub> films were synthesized. The study into the light transmission kinetics and characteristics of spectral electrochromic coloration revealed some differences electrochromic processes in these films. It was found that in the series of WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub>, CuWO<sub>4</sub>•WO<sub>3</sub> there is a slowing down of lithium intercalation in films. This is due to diffusion limitations in the process of coloring complex

oxides of Bi, Cu, and W. The spectral characteristics of electrochromic coloration Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> and CuWO<sub>4</sub>•WO<sub>3</sub> also differ from WO<sub>3</sub>, in that, in addition to tungsten trioxide, the contribution to light absorption is also made by oxides of Bi and Cu, which also partially are reduced by lithium in the process of their coloring. From this, it can be concluded that tungstates of metals can become effective electrochromic materials with an additional absorption band.

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# СРАВНИТЕЛЬНЫЙ АНАЛИЗ ЭЛЕКТРОХРОМНЫХ СВОЙСТВ ТОНКИХ ПЛЕНОК CuWO<sub>4</sub>•WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> И WO<sub>3</sub>

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**Аннотация:** Пленки  $Bi_2WO_6 \bullet WO_3$ ,  $CuWO_4 \bullet WO_3$  and  $WO_3$  были синтезированы комбинированным химическим и электрохимическим методами. Изучение кинетики светопропускания и спектральных характеристик электрохромного эффекта позволило установить некоторые различия электрохромных процессов в этих пленках. Показано, что в ряду  $WO_3$ ,  $Bi_2WO_6 \bullet WO_3$ ,  $CuWO_4 \bullet WO_3$  происходит замедление интеркаляции лития в пленки, что связано с диффузионными ограничениями в процессе окрашивания сложных оксидов Bi, Cu и W. Характеристики электрохромного окрашивания  $Bi_2WO_6 \bullet WO_3$  и  $CuWO_4 \bullet WO_3$  также отличаются от  $WO_3$  тем, что, помимо триоксида вольфрама, вклад в поглощение света вносят также оксиды Bi и Cu, которые также частично восстанавливаются литием в процессе их окраски. Сделан вывод о том, что вольфраматы металлов могут стать эффективными электрохромными материалами, имеющими дополнительную полосу поглощения в видимой области спектра.

Ключевые слова: электрохромизм, вольфраматы металлов, электрохромные композиты

### CuWO<sub>4</sub>•WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub> VƏ WO<sub>3</sub> NAZİK TƏBƏQƏLƏRİN ELEKTROXROM XÜSUSİYYƏTLƏRİNİN MÜQAYİSƏLİ TƏHLİLİ

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**Xülasə:** Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub>, CuWO<sub>4</sub>•WO<sub>3</sub> və WO<sub>3</sub> nazik təbəqələr kombinə edilmiş kimyəvi və elektrokimyəvi üsullarla sintez edilmişdir. İşığın ötürülməsinin kinetikasının və elektroxrom effektinin spektral xüsusiyyətlərinin öyrənilməsi bu plyonkalarda elektroxrom proseslərdə bəzi fərqlər müəyyən etməyə imkan vermişdir. Göstərilmişdir ki, WO<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>•WO<sub>3</sub>, CuWO<sub>4</sub>•WO<sub>3</sub> sırasında litiumun təbəqələrə interkalyasiyası ləngiyir, bu da Bi, Cu və W mürəkkəb oksidlərinin rənglənməsi prosesində diffuziya məhdudiyyətləri ilə əlaqələndirilir. Belə nəticəyə gəlinib ki, işıq spektrinin görünən hissəsində əlavə udma zolağın olması səbəbindən metal volframatlar effektiv elektroxrom materiallar ola bilərlər.

Açar sözlər: elektroxromizm, metal volframatlar, elektroxrom kompozitlər