Compounds of tellurium and selenium as binary and ternary semiconductors are involved in a wide range of applications [1-3]: solar cells, photoelectrochemical and thermoelectrical devices, micromechanical systems, optical filters, optical recording materials, superionic materials, as well as sensor and laser materials. As an example, bismuth telluroseleminides having Bi$_2$(Te,Se)$_3$ composition are recognized as thermoelectric materials with the best Seebeck coefficient at room temperature. Obviously, all these applications play an important role in a global sustainable development.

The electrochemical preparation of Bi$_2$(Te,Se)$_3$, semiconductor compounds [2-11] has attracted considerable interest because of the advantages of electrolysis procedure than other physical, chemical or metallurgical procedures. Historically, the electroplating of BiTeSe films, as well as preparation of ternary nanowires or nanotubes, has been carried out since 2003 [4,5] using aqueous baths. The cathodic process implies the presence in the bath of all three elements (introduced as dissolved precursors), but the weak solubility of bismuth and chalcogenide compounds (salts, oxides) imposes generally an acidic electrolyte. In most papers concerning BiTeSe electrodeposition nitric acid aqueous solution was used, even if sometimes other acid electrolytes [HCl, HClO$_4$ or H$_2$SO$_4$ solutions] were chosen. The recommended baths ([M HNO$_3$ solutions] contain generally 1-10mM concentrations of Bi$^{3+}$, Te$^{4+}$ (as HTeO$_2$$^{+}$) and Se$^{4+}$ (as HSeO$_2$$^{+}$) ionic species. The selection of deposition voltage and metallic support nature were performed usually by cyclic voltammetry tests.

In the recent years, because room temperature ionic liquids were increasingly applied for electrochemical purposes, the electrophoretic of tellurium and selenium compounds films has been demonstrated, for instance using 1-butyl-1-methylpyrrolidinium bis (trifluoromethyl-sulfonyl) amide or 1-ethyl-3-methylimidazolium tetra-fluoroborate / chloride) as more environmentally friendly media alternative [12-14]. The advantages of these novel baths include: extended electrochemical window; extremely low vapour pressures; significantly reduced hydrogen evolution as compared with the acidic aqueous baths conventionally employed; easy to achieve deposition of a desired composition of semiconductor deposit; lower electrical energy consumption comparing with aqueous solutions. The greater thermal stability and higher conductivity of the ionic liquid is useful to obtain crystalline semiconductor films through direct electrodeposition at higher temperatures without subsequent annealing.

We recently have shown the possibility of electrodeposition of Bi, Te and Se as elements or binary compounds using ionic liquids consisted in eutectic mixtures of choline chloride (2-hydroxy-ethyl-trimethyl ammonium chloride) with urea [15,16], malonic acid [17,18] or ethylene glycol [19]. In this paper, we present the cyclic voltammery and electrochemical impedance spectroscopy results regarding the electrodeposition at 60°C of BiTeSe films from three choline chloride (ChCl) based ionic liquids, i.e. mixtures of ChCl with ethylene glycol (EG), malonic acid (MA) or oxalic acid (OXA). These ionic liquids used in the present work were first developed by Abbott et al. [20] and consist of deep eutectic mixtures due to hydrogen bonding interactions between ChCl and chloride ions from ChCl.

The characterization of films grown on Cu substrate by long time electrolysis was made by corrosion tests. The stability of BiTeSe films against aggressive media, including marine environment, has relevance for thermoelectric and photoelectrochemical applications, improving the efficiency of thermoelectric devices or solar cells. In this respect, we mention that despite of some preliminary studies [18,21] the corrosion resistance of such semiconductor films has not been extensively studied.
Experimental part

The investigated ionic liquids as background electrolytes were prepared as mixtures of choline chloride (99%) with ethylene glycol (ChCl-EG 1:2 mole ratio), malonic acid (ChCl-MA 1:1 mole ratio) and oxalic acid dihydrate (ChCl-OxA 1:1 mole ratio), respectively, all reagents being purchased from Aldrich. The appropriate binary mixtures were heated at above 90°C for 30 min until homogeneous colourless ionic liquids are formed; then, the liquids were kept at room temperature. BiCl₃ or Bi₂O₃ (from Aldrich), and TeO₂ and SeO₂ (both from Alfa Aesar) were used as precursors for dissolved ionic species of bismuth, tellurium and selenium, respectively. The molarities were calculated using density values of 1.0963, 1.2065 and 1.2122 g cm⁻³ corresponding to ChCl-EG, ChCl-MA and ChCl-OxA mixtures, values determined in our laboratory at 60°C working temperature.

Cyclic voltammograms and impedance (EIS) spectra were recorded using a Pt plate (0.5 cm²) or a Cu disc (0.196 cm²) as working electrodes in order to evidence the cathodic and anodic processes in non-stirred baths. Pt plate as auxiliary electrode and Ag wire (immersed in the same ionic liquid) as reference electrode were also used in the electrochemical cell. The stability in 0.5 M NaCl aqueous solution of films electrodeposited on commercial Cu sheets was determined by recording potentiodynamic polarization (Tafel) curves and EIS spectra. The corrosion cell contained Cu plate covered with BiTeSe film (0.636 cm² exposed area), a Pt plate as auxiliary electrode and Ag/AgCl as reference electrode.

Zahner IM 6 potentiostat and BioLogic Sci. Instr. potentiostat, both with FRA, were used for cyclic voltammetry (1-100 mVs⁻¹ scan rate), potentiodynamic polarization (3 mVs⁻¹ scan rate) and electrochemical impedance spectroscopy (10 mV ac voltage, 200 kHz - 50 mHz frequency range).

Results and discussions

Regarding the relatively good solubility of bismuth, tellurium and selenium precursor compounds in ChCl-EG, ChCl-MA or ChCl-OxA ionic liquids in such amounts to have a significant electrochemical response for Bi³⁺, Te⁴⁺ and Se⁴⁺ ionic species, an assumption may be the presence of chloride anion Cl⁻ as ligand which forms complexes with these species; this complexation may improve the solubility of compounds and also the electrode potentials may close to each other in order to obtain a co-electrodeposition process. For instance, by dissolution of bismuth chloride (or bismuth oxide) precursor in the electrolyte we supposed the formation of BiCl₄⁻ complex species, leading to a lower diffusive ionic species than simpler Bi³⁺ ion [16,17]. Correspondingly, the tellurite (TeO₂⁻) or selenite (SeO₂⁻) dissolved precursors should be converted into their chloride complexes; alternatively, they may be present as acids (H₂TeO₃, H₂SeO₃) due to a significant presence of water in the ionic liquid, especially in ChCl-OxA mixture.

Cyclic voltammograms

The analysis by cyclic voltammetry (CV) technique was carried out to determine the appropriate potential range in which Bi³⁺, Te⁴⁺ and Se⁴⁺ cations co-deposit as BiTeSe films. We selected the CV curves obtained by sweeping the potential from anodic region towards cathodic direction (up to -1.2 V limit) and returning to anodic region (up to limit of +1.2 V) and back. It is important to mention that the CVs on Pt in all background ionic liquids recorded in an enlarged potential range, -1.4V ÷ +1.4V, did not show any electrode process.

Comparative cyclic voltammograms of single Te⁴⁺ and Se⁴⁺ ionic species and of ternary mixture of Bi³⁺, Te⁴⁺ and Se⁴⁺ ions recorded on Pt electrode in non-stirred choline chloride - ethylene glycol (1:2 mole ratio) ionic liquid are presented in figure 1.

It is clearly noticed that, obeying the standard electrode potential scale [1], selenium element deposits at the most positive potentials (plateau of a current in the potential range -0.2 ÷ -0.4 V, curve 1) and deposition of tellurium element is at more negative potentials (plateau around -0.6 V, curve 2). As an interpretation of the cathodic branch of voltammogram for BiTeSe ternary film deposition (curve 3), three reduction processes can be identified, with three distinct potential ranges of limited current or current peak. The first cathodic process that occurs on the curve 3 may be attributed to the Se⁴⁺ reduction, with the onset potential similar with curve 1 and a plateau in the potential range -0.1 ÷ -0.4 V). The second process may be attributed to co-deposition of ternary BiTeSe compound (with peak at -0.8 V). Finally, the third reduction process at potentials within -1.0 ÷ -1.2 V range may be due to deposition of a Bi-rich ternary BiTeSe layer. It results that Pt support first covers with a Se layer, and then the more cathodic polarization produces a co-deposition of Bi, Te and Se as ternary compound layer. An excessive cathodic polarization leads to a Bi massive deposition together with BiTeSe deposition and probably to the evolution of hydrogen or reduction of background electrolyte at over -1.2 V potential. Correspondingly, three distinguishable processes may be identified on the anodic branch of voltammogram: by scanning the potential towards positive values, the first two waves are attributed to anodic dissolutions of layers produced in the last reduction processes and the anodic peak (located at around +0.1 V) corresponds to the stripping of selenium layer previously formed on Pt.

Figure 2 compares the voltammetric response of the choline chloride – malonic acid (1:1 mole ratio) mixture in the presence of Bi³⁺ (as Bi₂O₃ in this system), Te⁴⁺ and Se⁴⁺ cations by varying the scan rate; the Pt electrode and working temperature were identical with binary system in figure 1. A narrow range of potentials and also low scan rates were selected for performing CVs in order to clearly show the existence of three couples of reduction/oxidation processes. All cathodic branches in this system have...
relatively similar shape as in figure 1, with plateaux of current (first from +0.15 to 0 V, the second one from -0.1 to -0.18 V) or a peak current (located in the range -0.2 ÷ -0.25 V).

We may suppose approximately a similar order of deposition: elemental Se layer, BiTeSe layer and Bi-rich BiTeSe layer. We observed that by increasing scan rate the last cathodic process is less evident, being overlapped with evolution of hydrogen (H+ ion coming from malonic acid dissociation). All currents on voltammograms in this system have lower values than in ChCl-EG, an explanation being the lower concentration of Bi3+, Te4+ and Se4+ ionic species. The difference in potential values comparing to figure 1 is certainly due to the shifting of potential for silver pseudoreference electrode by changing ionic liquid nature (malonic acid instead of ethylene glycol). On the anodic branches of voltammograms the first oxidation waves correspond to dissolution of Bi-rich ternary layer and BiTeSe layer, and the last anodic peak (in a sharp shape) represents a typical stripping process, which is consistent with the dissolution of electrodeposited selenium. On both branches of CVs in figure 2 all waves and peaks increase in height and remain at about the same potentials with various scan rates proving a diffusive control and relatively high reversibility of electrode processes.

Typical CVs recorded at various scan rates showing the electrochemical behaviour in conditions of simultaneous presence of Bi3+, Te4+ and Se4+ ions in choline chloride - oxalic acid (1:1 mole ratio) mixture are presented in figures 3 (a,b). It can be seen that there are some differences comparing to the previous figures, although the sequence of electrode processes seems to be similar. As Figure 3a shows, at lower scan rates (3-20 mVs-1) the Se deposition (represented by a plateau ranging from +0.4 to -0.1 V) is followed by a well defined peak (at -0.2 V) attributed to direct BiTeSe codeposition. However, the deposition of Bi in majority together with the ternary compound does not appear as distinct process because its wave is probably already overlapped with cathodic evolution of hydrogen (the continuous increase of current at over -0.25 V). This behavior may be explained by existence of significant concentration in H+ ions, provided from both oxalic acid dissociation and water coming as dihydrate. Figure 3b presents supplementary two CV curves recorded at higher scan rates (50 and 100 mVs-1). Now, the increase with scan rate of limiting current along the plateau of Se deposition is predominantly and, therefore, BiTeSe codeposition appears as a shoulder instead of a peak. Only two well contoured peaks are noticed on the anodic part of CVs, corresponding to consecutive dissolution of outer BiTeSe layer and of inner Se layer (as a sharp peak, typical for stripping). We notice that the other processes occurred in figures 3 to more positive potentials than +0.9 V are hardly to be explained, being certainly due to the content in oxalic acid and water of this ionic liquid system.

Interesting results were obtained during preliminary CV experiments in ChCl-OxA (1:1) ionic liquid by changing the nature of working electrode: Cu electrode was used instead of Pt working electrode. First it is worth to mention a displacement with ca. 600-700 mV more negatively of the stationary electrode potential. Figure 4 presents...
examples of cathodic branches of CV curves in ChCl-OxA ionic liquid on Cu electrode by keeping the same working conditions (temperature, scan rates) as in figures 3. It may be observed that the first process of Se deposition, that is located at most positive potential, is now almost hindered to be viewed; only it can be guessed this process as a small wave located around -0.27 V. So, we suppose that the main process of BiTeSe direct co-deposition occurs starting from -0.35 V and continues up to -0.5 V as a plateau (at low scan rates) or rather as a shoulder at higher scan rates. By polarizing more cathodically over -0.5 V, the continuous increase of current is responsible to the the third process discussed above (massive Bi deposition) that interferes with evolution of hydrogen. The anodic branches of CV curves show only a continuous increase of current, until around 0 V potential limit; we did not perform scanning at more positive potentials in order to prevent the electrochemical dissolution of copper support.

In a tentative to propose a mechanism of cathodic processes in two steps we may write the following equations:

\[ \text{Se}^{4+} + 4 \text{e}^- \rightarrow \text{Se} \text{ (film)} \]  
\[ 2 \text{Bi}^{3+} + (3-y) \text{Te}^{4+} + y \text{Se}^{4+} + 18 \text{e}^- \rightarrow \text{Bi}_2\text{Te}_3\text{ySe}_y \]  

In another way, we can take into consideration the equations proposed by all authors in aqueous solutions, where tellurium and selenium species were written as \( \text{H}_2\text{TeO}_3 \) and \( \text{H}_2\text{SeO}_3 \):

- for the first process (single deposition of Se with \( \text{H}^+ \) ions involvement):

\[ \text{H}_2\text{SeO}_3 + 4 \text{e}^- + 4 \text{H}^+ \rightarrow \text{Se} + 3 \text{H}_2\text{O} \]  

- for the second process:

\[ 2 \text{Bi}^{3+} + (3-y) \text{H}_2\text{TeO}_3 + y \text{Se}^{4+} + (12-4y) \text{H}^+ + 18 \text{e}^- \rightarrow \]  
\[ \text{Bi}_2\text{Te}_3\text{ySe}_y + (9-3y) \text{H}_2\text{O} \]

Of course, the massive deposition of Bi means a direct reduction with three electrons of \( \text{Bi}^{3+} \) ion as a simple electrode process.

In general, in spite of the obvious differences in peak potential separation and peak shapes we can appreciate that all the reduction/oxidation processes are diffusion controlled and moderately irreversible, and this irreversibility degree may be due to a large uncompensated ohmic drop inside the ionic liquid.

**Electrochemical impedance spectra**

The cathodic formation of films was also studied by the electrochemical impedance spectroscopy (EIS) measurements at 60°C. Figures 5 a,b show comparatively the recorded Nyquist and Bode (both impedance modulus and phase angle vs. frequency dependences) diagrams recorded on Pt in choline chloride – ethylene glycol ionic liquid. The electrode was polarized in the potential region where the reduction process has a significant cathodic current (peak or limiting current on CV curves). For tellurium deposition (curve 1) the Nyquist spectrum consists of a capacitive loop, followed by a straightline of unit slope. This shape clearly indicates a formation of Te film onto the Pt electrode surface. Curve 2 in the same figure represents two consecutive semicircles followed also by a straightline of unit slope and may be interpreted as a process where the first step is Se element deposition and the second step is a codeposition of TeSe semiconductor compound which has a composition of solid solution. Curve 3 corresponds to ternary codeposition resulting BiTeSe compound; the diameter of this semicircle (which represents the transfer charge resistance, \( R_{ct} \)) is now very small indicating a higher current of the faradaic process. It can be seen from figure 5b that according to the phase angle values, TeSe semiconductor has the highest insulating properties (-60° phase angle) due to its high content of selenium, Te film is also a good semiconductor material (-55°), but the obtained BiTeSe film has already a metallic character (-30° phase angle), explained by a significant content of Bi and low content in Se in the ternary compound.
By gradually polarizing Pt electrode in negative direction, first successive Nyquist and Bode diagrams (fig. 7 a,b) suggest the existence of two consecutive deposition processes, as two semicircles or two phase angle maxima in the potential region where deposition of both Se and BiTeSe takes place. For more negative potentials than -0.25 V, the deposition of ternary BiTeSe film on already deposited selenium becomes predominantly, so a single Nyquist semicircle and a single phase angle maximum is recorded. Normally, the magnitude of semicircle diameters are different, because the diameter decreases with cathodic polarization indicating an increase of charge transfer rate and faradaic current during deposition. All Nyquist diagrams do not show perfect semicircles, rather depressed semicircles, and this fact has been attributed to frequency dispersion due to inhomogeneity and roughness of electrode surface.
The EIS diagrams obtained for Cu electrode (fig. 8) confirm the interpretation of cyclic voltammetry curves recorded using this electrode, respectively a quite direct BiTeSe deposition followed by massive deposition of Bi at very negative potentials. Thus, the successive Nyquist single semicircles (fig. 8a) decrease gradually in their diameter with cathodic polarization and the successive Bode curves (fig. 8b) decrease dramatically their values of phase angle maxima, reaching even to -20° for -0.50 V polarization, that is a phase angle value characteristic for a metallic film.

We can validate and quantitatively interpret the experimental impedances obtained, by modeling the electrode process using an equivalent electrical circuit. Figure 9 shows an equivalent circuit established using the specialized software Zview 2.90c (Scribner Assoc.) and employed to fit the EIS experimental data. Regarding this circuit, the whole impedance of the electrode is represented as a series connection of the solution resistance Rs and a parallel circuit of the double-layer capacitance (which is generally denoted as Cdl) and charge transfer resistance, Rct, here denoted as R1. However, we showed that the electrode surface is modified by microscopic roughness caused by scratches, pits, etc., and also its capacitance is dispersed, being influenced by slow adsorption of ions and chemical inhomogeneities of the surface. In such cases a constant phase element CPE is often introduced instead of Cdl. Its impedance is given by:

\[ Z_{\text{CPE}} = \frac{1}{T(j\omega)^P} \]

where \( T \) is a constant in \( \text{F cm}^{-2} \), \( \omega \) is angular frequency \( (\omega = 2\pi f, f - \text{ac frequency, Hz}) \) and \( P \) is an exponent. Only when \( P = 1 \), \( T = C \), and purely capacitive behavior is obtained. In general, eq. (5) may represent pure capacitance for \( P = 1 \), infinite Warburg impedance for \( P = 0.5 \), pure ohmic resistance for \( P = 0 \) and pure inductance for \( P = -1 \). W1 is a Warburg element of linear finite-length diffusion.

Table 1 contains the values of equivalent circuit elements for the best fitting of experimental data representing BiTeSe deposition in choline chloride + oxalic acid (1:1) ionic liquid, 60°C. It may be seen that the changes in EIS spectra after polarization of Pt or Cu electrode at different cathodic overpotentials were well reflected in the fitted parameters.

![Fig. 9 Scheme of equivalent electrical circuit for BiTeSe films obtained from choline chloride + oxalic acid (1:1) ionic liquid, 60°C. Circuit components: Rs – solution ohmic resistance; CPE 1 – constant phase element replacing double layer capacitance; R1 – charge transfer resistance (Rct for cathodic deposition) or polarization resistance (Rp for corrosion tests); W – Warburg diffusion impedance](http://www.revistadechimie.ro)

**Preparation and corrosion behaviour of BiTeSe films deposited on Cu support**

BiTeSe film samples were prepared under potentiostatic or galvanostatic control on copper sheets from mixtures of choline chloride with malonic acid or with oxalic acid at 60°C, without stirring the ionic liquid bath. Table 3 presents the operating conditions in both ionic liquid electrolytes and some characteristics of BiTeSe deposits (aspect, adherence on copper support).

We consider that deposits formed in shorter time (BiTeSe-1 and BiTeSe-3 samples) have a composition with relatively higher Se content than the other deposits. We noticed that the BiTeSe films electrodedeposited in conditions of long-time electrolysis (60 or 120 min) possess a smoother surface and more compact structure than those prepared by the short-time electrolysis. It is also confirmed (from color changes) that electrodeposition at more negative potentials leads to an increase of bismuth content in the BiTeSe film. So, it is clear that the characteristics of...
deposits may be easily controlled by selecting the appropriate electrolysis conditions. Samples elaborated under potentiostatic (denoted as 1,2) or galvanostatic (denoted as 3,4) control using bath containing choline chloride - malonic acid (1:1) mixture +2.5 mM Bi^{3+} + 2.5 mM Te^{4+} + 0.5 mM Se^{4+} were tested for their corrosion resistance. Figures 10 a,b show the potentiodynamic polarization curves recorded for these samples. Also, a comparison of electrochemical impedance spectra (EIS) recorded by polarizing BiTeSe films at different anodic overpotentials can be seen in figures 11-14. We mention that more corrosion data regarding our investigations in various liquid media (including ionic liquids) were reported in another paper [18].

In all cases presented in figures 10 the Tafel polarization curves were plotted starting from the stationary potential firstly toward cathodic direction and then toward anodic direction. The increase in current in the first part of anodic branch starting from stationary potential is indicative of the active dissolution of BiTeSe film. However, at a potential around +0.1 V vs Ag/AgCl reference electrode a decrease in the current for all samples may suggests a passivation process that may be due to the insoluble corrosion products formed and adsorbed temporary on the surface. Then, in more positive part of Tafel curve the film returns to an active state and continues to be corroded with a quite moderate constant rate; only samples with more Se content (BiTeSe-1 and BiTeSe-3) show a new passivation process at high anodic polarization. We consider that the cathodic process when the applied potential increased into the negative region (cathodic branch of Tafel curves) corresponds mainly to the evolution of the hydrogen that is an activation-controlled process.

Estimation of the corrosion parameters was made using the corrosion soft of potentiostats, the characteristic corrosion data being listed in table 4. It seems, from corrosion currents values, that more corrosion resistance is reached by galvanostatic electrolysis; (samples 3 and 4); also, in these cases the cathodic (bc) and anodic (ba) Tafel slopes were obtained in a narrow range (83-94 mV/decade).

Also, a comparison of electrochemical impedance spectra (EIS data) recorded by polarizing BiTeSe films in 0.5 M NaCl solution at different anodic overpotentials exhibited a quite similar corrosion behaviour (figs. 11-14).

### Table 2
VALUES OF COMPONENTS IN THE EQUIVALENT CIRCUIT USED FOR BEST FITTING OF EIS DATA FOR BiTeSe DEPOSITION AT VARIOUS CATHODIC POTENTIALS ON Cu ELECTRODE. IONIC LIQUID: ChCl-OxA (1:1) + 10 mM BiCl_{3} + 5 mM TeO_{2} + 5 mM SeO_{2}, 60 °C

<table>
<thead>
<tr>
<th>Component in scheme of Fig. 9</th>
<th>Values of components in the equivalent circuit for various cathodic potentials</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-0.35 V</td>
</tr>
<tr>
<td>$R_{c}$, $\Omega$</td>
<td>22.7</td>
</tr>
<tr>
<td>CPE1-T $\times 10^{6}$</td>
<td>50</td>
</tr>
<tr>
<td>CPE1-P exponent</td>
<td>0.86</td>
</tr>
<tr>
<td>$R_{1}$, $R_{2}$, $\Omega$</td>
<td>779</td>
</tr>
<tr>
<td>W1-I, $\Omega$</td>
<td>1958</td>
</tr>
<tr>
<td>W1-T (pseudoinductance), $H$</td>
<td>1.06</td>
</tr>
<tr>
<td>W1-P exponent</td>
<td>0.31</td>
</tr>
</tbody>
</table>

### Table 3
OPERATION CONDITIONS DURING PREPARATION OF BiTeSe SAMPLES ON COPPER SHEET USING ChCl-MA AND ChCl-OxA IONIC LIQUIDS, 60 °C

<table>
<thead>
<tr>
<th>Bath</th>
<th>Sample</th>
<th>Controlled current or potential</th>
<th>Electrolysis time, min</th>
<th>Characteristics of deposit</th>
</tr>
</thead>
<tbody>
<tr>
<td>ChCl-MA (1:1)</td>
<td>BiTeSe-1</td>
<td>-0.55 V vs. Ag ref.</td>
<td>5</td>
<td>Uniform, light grey, adherent</td>
</tr>
<tr>
<td>+2.5 mM Bi^{3+} + 2.5 mM Te^{4+} + 0.5 mM Se^{4+}</td>
<td>BiTeSe-2</td>
<td>-0.55 V vs. Ag ref.</td>
<td>10</td>
<td>Uniform, grey, adherent</td>
</tr>
<tr>
<td>ChCl-OxA (1:1)</td>
<td>BiTeSe-3</td>
<td>2.5 mA/cm²</td>
<td>10</td>
<td>Uniform, grey-black, adherent</td>
</tr>
<tr>
<td>+10 mM Bi^{3+} + 5 mM Te^{4+} + 5 mM Se^{4+}</td>
<td>BiTeSe-4</td>
<td>3 mA/cm²</td>
<td>60</td>
<td>Uniform, compact bright grey-black, adherent</td>
</tr>
<tr>
<td>ChCl-OxA (1:1)</td>
<td>BiTeSe-I</td>
<td>-0.35 V vs. Ag ref.</td>
<td>120</td>
<td>Uniform, bright black, adherent</td>
</tr>
<tr>
<td>+10 mM Bi^{3+} + 5 mM Te^{4+} + 5 mM Se^{4+}</td>
<td>BiTeSe-II</td>
<td>-0.42 V vs. Ag ref.</td>
<td>60</td>
<td>Uniform, compact, bright black, adherent</td>
</tr>
</tbody>
</table>

Fig. 10. Semilogarithmic potentiodynamic polarization plots for BiTeSe samples 1,2,3,4 (table 3) in 0.5 M NaCl aqueous solution; 3 mVs⁻¹ scan rate, 25 °C, 0.636 cm² exposed area
Nyquist plots are generally in a shape of capacitive arc; the diameter of semicircles is gradually decreased by polarizing anodically, proving decreased polarization resistance, \( R_p \), i.e. an increase of corrosion current, as expected. However, the shape of impedance spectra is not ideal semicircle. We consider that the factors such as surface roughness, the existence of corrosion products or formation of passive films can contribute to a deviation from ideal semicircle.

The behaviour of deposit corrosion was simulated with some equivalent electric circuits as models; one was shown in figure 9 and the other two models are presented in figures 15. The circuit in figure 15a is a simple Randles
Table 5
FITTING PARAMETERS FOR BiTeSe-1 SAMPLE ANODICALLY POLARIZED IN 0.5 M NaCl AQUEOUS SOLUTION USING EIS TECHNIQUE, 25 °C. EXPOSED SURFACE: 0.636 cm²

Table 6
FITTING PARAMETERS FOR BiTeSe-2 SAMPLE ANODICALLY POLARIZED IN 0.5 M NaCl AQUEOUS SOLUTION USING EIS TECHNIQUE, 25 °C. EXPOSED SURFACE: 0.636 cm²

Table 7
FITTING PARAMETERS FOR BiTeSe-3 SAMPLE ANODICALLY POLARIZED IN 0.5 M NaCl AQUEOUS SOLUTION USING EIS TECHNIQUE, 25 °C. EXPOSED SURFACE: 0.636 cm²

Table 8
FITTING PARAMETERS FOR BiTeSe-4 SAMPLE ANODICALLY POLARIZED IN 0.5 M NaCl AQUEOUS SOLUTION USING EIS TECHNIQUE, 25 °C. EXPOSED SURFACE: 0.636 cm²

circuit with a pure capacitor having a double-layer capacitance (Cdl). The second one model (fig. 15b) contains a scheme where the connection of the polarization resistance (denoted as R₁) and W₁ in parallel with a constant phase element (CPE₁) is in series with a second parallel connection of BiTeSe film components: ohmic resistance (R₂) and its capacitance (CPE₂).

Tables 5-8 contains the fitting data using circuit models from Figures 9 and 15a. Although we did not perform calculations with with circuit model from figure 15b, we consider that this last scheme is better suitable for interpretation of several EIS spectra showing two time constants (two semicircles or two phase angle maxima). The obtained data demonstrate clearly that BiTeSe films formed by long-time electrolysis (BiTeSe-2 and BiTeSe-4) have a higher corrosion resistance (they have higher Rp values) than films formed in short time. The highest metallic character seems to be for BiTeSe-4 proving that it has a composition with higher Bi content.

Conclusions
The present electrochemical investigation showed that BiTeSe films can be successfully deposited from all three ionic liquids electrolytes consisting in binary mixtures of choline chloride with ethylene glycol, malonic acid and oxalic acid, respectively. From cyclic voltammetry and impedance experiments carried out at 60 °C using Pt and...
Cu electrodes there were evidenced the characteristics of cathodic processes and, especially, it was possible to predict the potential ranges for codeposition of Bi, Te and Se as ternary BiTeSe compound.

The electrodeposition of BiTeSe films takes place from all electrolytes on a Se-covered substrate. Also, these films were formed at less negative potentials than for deposition of singular Se (or Te films). In general, the corrosion resistance of BiTeSe films in 0.5 M NaCl solution was similar with other semiconductor films containing Se or Te and in the evolution of corrosion may possible occur retaining of corrosion products and formation of a passive film.

Finally, we mention that the preparation of BiTeSe films by electrodeposition technique from choline chloride – oxalic acid ionic liquid was reported for the first time in literature.

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References
1. BOUROUSHIAN, M., Electrochemistry of Metal Chalcogenides, Springer-Verlag, Berlin, 2010, p. 60
3. BOULANGER, C., J. ELECTRON. MATER., 39 (9), 2010, p. 1818

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