Higher polythionic acids – the precursors for formation of thallium sulphide layers on polyethylene surface

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The method of thallium sulphide thin films deposition on low density polyethylene (LDPE), based on applying solutions of higher polythionic acids (H$_2$S$_n$O$_{6n}$, $n = 21, 33, 45$), was developed and investigated. The concentration of sulphur in the LDPE film increases with the increase of temperature of higher polythionic acid solution and its sulphurity, as well as the duration of retention of LDPE in its solution is prolonged. The sulphur concentration in the PE film under the same duration of sulphurisation and repetition of treatment cycles insignificantly changes. When the sulphurised LDPE was treated with a thallium Tl(I) salt solution, sulphur present in the polymer reacts with thallium ions and the layers of thallium sulphides, Tl$_x$S$_y$, of different composition are formed. The stoichiometric composition of the layer changes from Tl$_{0.1}$S to Tl$_{2.6}$S. The amount of thallium increases in the sulphide layers when the number of the cycles of LDPE sulphurisation in the H$_2$S$_{33}$O$_{6}$ solution and treatment of sulphurised polymer with the Tl(I) salt solution is increased. The stoichiometric composition of Tl$_x$S$_y$ coats of one treatment cycle samples changes from Tl$_{0.5}$S to Tl$_{1.1}$S. The X-ray diffraction analysis of the polyethylene, treated with the solution of higher polythionic acid, H$_2$S$_{33}$O$_{6}$, and then with the Tl(I) salt solution, confirmed the premise that on the LDPE surface the layers of thallium sulphides, Tl$_x$S$_y$, were formed. Five phases of thallium sulphides were identified: TlS, Tl$_2$S, Tl$_3$S$_2$, Tl$_4$S$_3$, and Tl$_5$S$_5$. The TlS phase prevailed in diffractograms. The chemical and phase composition of the Tl$_x$S$_y$ layers surface (up to 1 nm depth) is identified by X-ray photo-electronic spectroscopy analysis. The analysis data confirmed that the compounds of TlS, Tl$_2$S, Tl$_2$SO$_4$, Tl(OH)$_3$, S$_8$ and Tl$_2$O$_3$ were formed on the surface of the layers. The morphology of thallium sulphide layers and the mechanism of their formation are determined by means of scanning atomic force and electronic microscopy. Two processes take place during the formation of the layer: desorption of sulphur and reaction of thallium ions from the thallium(I) salt solution with sulphur present on the LDPE surface. The layer of thallium sulphides is formed on the surface of the sulphur layer. Deposition of thallium sulphide films from the solution of higher polythionic acid on the LDPE films is not homogeneous, it has the separated island type morphology. The identified regularities let purposefully choose the conditions for sulphurisation of LDPE films by using the solution of higher polythionic acid as the agent of polymer sulphurisation in order to compose the layers of thallium sulphides, Tl$_x$S$_y$, of the desired chemical and phasic composition on the surface of this polymer by means of the sorption–diffusion method.

Keywords: polyethylene, thallium sulphide thin films, atomic force microscopy, X-ray diffraction, X-ray photoelectron spectroscopy
INTRODUCTION

Composite materials (composites) are distinguished by the properties of their components and are needed for advanced technologies. Modification of elastic, light and resistant to the environment polymeric materials by creating thin layers of compounds with important physical properties on their surface leads to the formation of materials with new properties [1-5]. These thin layers have a number of applications in various fields, including coatings, interference filters, polarizers, narrow band filters, solar cells, photoconductors, IR detectors, waveguide coatings, magnetic and superconducting films, microelectronic devices, etc. [6-8]. Many metal sulphide compounds have excellent optical properties in the visible and IR region of the spectrum [10-17].

Recently at the Department of Inorganic Chemistry the sorption–diffusion method of low density polyethylene (LDPE) modification by semiconductor and electrically conductive layers of CuS has been investigated. By means of this method, first of all, PE is saturated with sulphur treating with the solutions of higher polythionic acids, HnS2O (n = 9–45) [8, 9], and then it is treated with a water solution of copper (II) salts [13, 14]. The layers of thallium sulphides are distinguished by a great variety of stoichiometry and valuable physical properties. The layers of thallium sulphides are commonly produced by the deposition method from solutions [15-17]. But its formation is more convenient by the use of sorption methods.

The aim of the present work was to summarize and to discuss the results obtained by us studying the formation and characterization of thallium sulphide layers on LDPE films surface by the sorption–diffusion method, since mainly only separate fragments of these studies in our previous publications have been described [18-45].

EXPERIMENTAL

The layer of thallium sulphide was deposited on the films of low density polyethylene (LDPE) (GOST 10354-82), produced at Vilnius Plastic Plant ‘Plasta’. The thickness of the LDPE film was 0.22±0.01 mm. Before experimenta- tion the 15 × 75 mm samples of LDPE were accordingly prepared. They were washed for 10 min by a 4% solution of a cationic surfactant, degreased in benzene for 10 min. The extracted samples were washed in distilled water and dried. Distilled water, reagents of the grades ‘especially pure’, ‘chemically pure’ and ‘analytically pure’ were used in the experiments.

The samples of LDPE were sulphured in 0.002 mol·dm−3 stirred in the solutions of higher polythionic acids (HnS2O, n = 21, 33, 45) at 25, 40, 60 and 80°C. Then these solutions were kept in the darkness in an open vessel for 48 h until hydrogen sulphide removed fully [51].

The solution of thallium(I) salt was made from crystalline thallium(I) sulphate. The concentration of thallium(I) sulphate solutions was 0.01, 0.05 and 0.1 mol·dm−3 (pH 2.8) with addition of 0.015 mol·dm−3 KOH (pH ~ 11.4). When using alkaliified thallium(I) salt, thallium sulphide layers on the surface of LDPE were obtained [40, 45]. For the formation of TlS films, the sulphured LDPE samples were treated with the Tl(I) salt solution at 80°C.

The concentration of diffused sulphur into LDPE was determined by several methods. The sulfur was extracted with acetone and the content in the extract was determined by the spectrometric cyanide method using a Specord UV/VIS spectrophotometer (λ = 450 nm) [52]. The concentration of sulphur with more thick TlS layers, which have been obtained repeating 3 times the sulphurisation of LDPE, was determined by the nephelometric method, using an atomic absorption spectrophotometer Perkin-Elmer 503 (λ = 553.6 nm), indirectly by the measurements of Ba2+ ions after sulphur deposition in the form of BaSO4 [53].

The amount of Ti in LDPE was determined using an atomic absorption spectrophotometer Perkin-Elmer 503 (λ = 276.5 nm) [53].

The phase composition of thallium sulphides layers on the surface of LDPE was investigated by means of X-ray diffraction (XRD) with a DRON-6 diffractometer provided with a special device for beam limitation at low and medium diffraction angles using a graphite-monochromatized Cu-Kα radiation source (λ = 1.54178 Å) under 30 kV voltage and 30 μA current. The XRD patterns were recorded with a step size of 0.05° from 20 = 10° to 70°. X-ray diffractograms of LDPE samples with the layers of thallium sulphides were treated using the program Search Match to eliminate the maxima of LDPE. The experimental XRD patterns with the layers of TlS analysis were performed using the Crystallografica software and PDF data base [17, 37, 54, 55].

The chemical composition of thallium sulphides layers was investigated by the method of X-ray photoelectron spectroscopy (XPS). Spectra were recorded by means of a spectrometer ESCALAB MKII, produced at VG Scientific (England), the power of which is 300 W, applying the radiation of Mg anode (Mg Ka 1253.6 eV, capacity 300 W). In the analytic chamber the vacuum of 1.33·10−9 Pa was maintained. In order to identify the distribution of elements by depth, the signal of Ar+ ions, the ions flow energy of which is approximately 1.0 keV, was applied. The samples were etched in a preparatory chamber under 9.31·10−3 Pa vacuum and 20 μA current, this corresponds to 20 ā/min etching speed. The etching time was 10 s. To investigate the layers obtained by the XPS method the photoelectron spectra of Ti 4f, O 1s and S 2p were recorded. Empirical sensitivity factors for these elements were taken from the literature [56, 58] and the spectra obtained were compared with the standard ones [59].

The morphology of the surface of TlS layers was inves- tigated by atomic force microscopy (NT-MDT Inc.,
Zelenograd, Moscow, Russia) in the contact mode using commercial (Nano Technology Instruments – Europe BV) Si cantilevers CSG10 series with 0.2 Nm⁻¹ force constant and 10 nm tip curvature. The data of measurements were analysed by means of the programs Scan Atomic™, SPIP.

RESULTS AND DISCUSSION

The preparation of layers of thallium sulphides by sorption–diffusion on the surface of LDPE

The formation of thallium sulphide, Tl₂S₃, layers on the LDPE surface consists of two parts. During the first part of the process the samples of LDPE were sulphured in a water solution of higher polythionic acids H₂S₉O₅ (n = 21, 33, 45) as a chalcogenization agent. The molecules of elemental sulphur are formed during the decomposition of higher polythionic acids and are then sorbed-diffused into the matrix of the LDPE surface.

During the second part of this process the sulphured samples are treated with a water solution of Tl(I) salt. Thallium sulphide layers on the LDPE surface are formed by the sorption–diffusion method as a result of the oxidation–reduction reaction between sulphur firstly sorbed-diffused in LDPE and Tl⁺ ions present in a solution of thallium(I) salt. Two processes take place during the formation of the thallium sulphide film on the surface of LDPE: elemental sulphur desorption and interaction of Tl⁺ with elemental sulphur in the solution.

Sulphurisation of polyethylene by solutions of higher polythionic acids H₂S₉O₅

The samples of the LDPE film were sulphurised in solutions of higher H₂S₉O₅ (n = 21, 33, 45) at 60°C temperature. It was known that the molecules of elemental sulphur are formed during the decomposition of higher polythionic acids. The concentration of sulfur adsorbed-diffused into LDPE depends on the degree of acid sulphurity (n), the temperature of the solution and the period of polymer treatment.

The concentration of sulphur sorbed-diffused into LDPE has the increase of the number of sulphur atoms in the molecule of H₂S₉O₅ solutions. Further on prolonging the duration of sulphurisation, the sulphur concentration increased more slowly in LDPE, however, the saturated concentration was inaccessible even during 3 h (Fig. 1).

Thus, these results reveal that the solutions of higher polythionic acid H₂S₉O₅ are suitable to be applied for LDPE sulphurisation during further research, as a further increase of the sulphurisation degree (the number of S atoms in a molecule, n) does not make a more distinguished influence upon the effectiveness of LDPE sulphurisation. Figure 1 presents the change of sulphur concentration in the LDPE film when the samples are sulphured in the H₂S₉O₅ acid solution during different periods of time at 25, 40, 60 and 80°C temperature.

The kinetic curves of sulphur concentration changes show that the concentration of sulphur sorbed-diffused
into LDPE increases with the sulphurisation time at all temperatures used. But the rise of the temperature from 40 to 60°C makes a vivid influence on the values of sulphur concentration, \( c_s \), achieved in the polymer films. The saturated sulphur concentration in LDPE at 25°C temperature was reached after 1 h and at 40°C temperature it was reached after 2 h of treatment. While the samples of LDPE were treated in the \( \text{H}_3\text{S}_8\text{O}_6 \) solution at 60 and 80°C temperature, the saturated \( c_s \) was reached after ~2.5 h.

Thus our experimentations showed that the LDPE sulphurisation was purposive to be performed in the \( \text{H}_3\text{S}_8\text{O}_6 \) solution at 60°C temperature during further research.

Formation of thallium sulphide layers on the sulphurised surface of LDPE by applying thallium(I) salt solutions

The second part of LDPE modification is the reaction of a sulphurised film with a solution of TI(I) sulphate and formation of TI\(_3\)S\(_2\) layers on the LDPE surface.

The TI\(_3\)S\(_2\) layer on the surface of the LDPE film is formed when TI\(^+\) ions react with sorbed-diffused sulphur in the polymer. Thallium sulphide layers are formed in the surface of LDPE as a result of the oxidation–reduction reaction between sulphur sorbed by LDPE and thallium(I) ions present in the solution of thallium(I) salt [43]:

\[
3\text{Tl}^+ + 1/8\text{S}_8 \rightarrow \text{Tl}_2\text{S}_3 + \times\text{Tl}^{15+}.
\]

TI\(^+\) ions have higher mobility than S\(_8\) molecules, and this TI\(_3\)S\(_2\) layer is formed when TI\(^+\) ions diffuse into the sulphurised LDPE. Low-density polyethylene consists of unevenly distributed amorphous and crystal areas. Sorption–diffusion of sulphur as well as of TI\(^+\) ions is possible only in amorphous areas of LDPE. The product of the reaction impedes further diffusion of TI\(^+\) ions into the depth of the polymer, but there are enough defects of the structure in LDPE, where diffusion of small molecule compounds occurs even through the formed layer of TI\(_3\)S\(_2\). By the changed colour of the LDPE film it is possible to judge that, after processing the sulphurised LDPE with the TI(I) solution, the layer of thallium sulphides is formed on its surface [43]. The colour of LDPE films directly depends on the conditions of sulphurisation. The LDPE films, whose concentration of diffused sulphur was small, become redly brownish after ‘thalliumisation’, and the films, whose concentration of sulphur was larger, after the reaction with the TI(I) solution become dust-coloured or even blackish.

The amount of thallium in the polymer is influenced by the processes of LDPE sulphurisation and treatment with the TI\(_3\)SO\(_4\) salt solution.

It was found that the amount of thallium in the sulphurised layers on the LDPE surface increased with the prolongation of LDPE sulphurisation in the \( \text{H}_3\text{S}_8\text{O}_6 \) solution at all used temperatures. A different amount of thallium was found in the TI\(_3\)S\(_2\) layers on the surface of LDPE at every temperature [43]. The maximum value of the TI amount depends on the temperature of the sulphurisation solution (Fig. 3).

The amount of thallium directly depends on the sulphur concentration in LDPE, i.e. when the latter increases, the amount of TI also increases in the obtained layer on the LDPE surface (Fig. 3). This is quite understandable because if the concentration of sorbed-diffused sulphur is higher, a greater amount of TI\(^+\) ions can take part in the oxidation–reduction reaction with sulphur molecules.

The influence of experimental conditions (temperature, concentration and duration of LDPE sulphurisation and the treatment of sulphurised LDPE with thallium(I) salt solution) on the thallium sulphide layers in the LDPE surface was studied. The obtained results show that \( m_{Tl} \) constantly increases when the temperature of ‘thalliumisation’ is increased (Figs. 3, 4). The maximum amounts of thallium are obtained when sulphurised LDPE is ‘thalliumised’ in the solution of thallium (I) salt at 80°C temperature. The amount

Fig. 3. Changes of the thallium concentration in the TI\(_3\)S\(_2\) layer. LDPE was sulphurised in the \( \text{H}_3\text{S}_8\text{O}_6 \) solution, then treated with the 0.1 mol·dm\(^{-3}\) solution of TI\(_3\)SO\(_4\) at 80°C temperature for 10 min. The temperature of the sulphurisation solution, °C: 25 (1), 40 (2), 60 (3), 80 (4)

Fig. 4. Changes of the amount of thallium in the TI\(_3\)S\(_2\) layer on the sulphur concentration in the LDPE film. LDPE was sulphurised for 10 min, then treated with the 0.1 mol·dm\(^{-3}\) TI\(_3\)SO\(_4\) solution at 80°C temperature. The temperature of the \( \text{H}_3\text{S}_8\text{O}_6 \) solution, °C: 60 (1) and 80 (2)
of thallium in the TlS$_y$ layer increases unevenly as it depends on the temperature of LDPE sulphurisation. When the LDPE samples are sulphurised at 60°C temperature, $m_{Tl}$ hardly depends on the concentration of the thallium(I) salt solution (Fig. 5).

When the LDPE film samples are sulphurised at 80°C temperature, the concentration of the TlS$_{y}$ solution more influences the thallium amount in the TlS$_{y}$ layers (Fig. 5).

If we prolong the 'haliuminisation' of the sulphurised LDPE film with the thallium(I) salt solution, the amount of thallium in the TlS$_{y}$ layers will increase. Changes of the amount of thallium in the TlS$_{y}$ layers are shown in Fig. 6.

The prolongation of sulphurisation and the repetition of treatment cycles influences the amount of thallium in the TlS$_{y}$ layers quite differently.

Considering that the cycles of the process of TlS$_{y}$ layers formation can give a positive result, for example, to obtain thicker sulphide layers better cohered with polymer, we further investigated how concentrations of sulphur and thallium in LDPE change after each cycle. Thicker layers were obtained by repeating the cycles of sulphurisation and treatment with the thallium(I) salt solution for three times, i.e. sulphurisation-'thalliumisation', then sulphurisation of the same sample of the PE film and 'thalliumisation', etc.

The solutions of higher polythionic acid H$_2$S$_{33}$O$_6$ at 60°C temperature for this experimentation were used. The presented data show that the sulphur concentrations in the LDPE film, which was kept in H$_2$S$_{33}$O$_6$ solutions for different time intervals, increase as well. During the first cycle, in 15 min from the experiment beginning, the sulphur concentration equals $c_s$ ~ 1.95 mg·cm$^{-2}$, after 2 h the S concentration increases ~4 times and is equal to $c_s$ ~ 7.65 mg·cm$^{-2}$. By repetition of sulphurisation cycles, in the LDPE samples of two cycles of the treatment, sulphurised for 15 min, $c_s$ is equal to ~2 mg·cm$^{-2}$, it is 1.5 times larger after 30 min and ~2.5 times larger after 60 min. The sulphur concentration of three treatment cycles as well as one cycle in the LDPE samples of sulphurisation for 120 min is four times larger than in the sulphurised samples for 15 min (Fig. 5). Thus the sulphurisation duration makes a great influence upon the sulphur concentration in the LDPE film.

It is supposed that by repeating the cycles a moderate part of S$_y$ segregated in the solution of polythionic acid absorbs on the coat surface or reacts with the composed layer of thallium sulphide:
The results of the research permit to state that compounds possessing thallium and sulphur, i.e. layers of thallium sulphides, are formed on the surface of the PE film, first affected with solutions of higher polythionic acid $H_{2}S_{3}O_{6}$, and later with that of $\text{Tl(I)}$ salt.

The morphology of $\text{Tl}_x\text{S}_y$ layers on the surface of LDPE

To characterize the morphology of $\text{Tl}_x\text{S}_y$ layers on the surface of LDPE, the method of atomic force microscopy (AFM) was applied. Fields of $30 \times 30$ microns were investigated; for the quantitative estimation of the surface, the standard programs of the view treatment were used. Figures 8 shows the typical surface morphology of the deposited sulphur and thallium sulphide thin films on the LDPE surface obtained by AFM.

\[
\frac{(y-x)}{8}S_{y} + x\text{Tl}_x\text{S}_y \rightarrow y\text{Tl}_x\text{S}_y. \tag{2}
\]

Fig. 8. Changes of the sulphur concentration in LDPE on the number of cycles of its treatment in the $H_{2}S_{3}O_{6}$ solution. The sulphurisation duration of LDPE, min: 15 (1), 30 (2), 60 (3) and 120 (4)

Fig. 9. The study of the morphology of $\text{Tl}_x\text{S}_y$ layers on the LDPE surface by AFM: LDPE sulphurised during 12 min in the solution of $H_{2}S_{3}O_{6}$ at 60°C temperature (a); sulphurised LDPE treated with the 0.1 mol·dm$^{-3}$ solution of $\text{Tl}_2\text{SO}_4$ at 80°C temperature during 1 min (b)
Two processes take place during the formation of the layer: desorption of sulphur and reaction of thallium ions from the thallium(I) salt solution with the sulphur present on the LDPE surface. The layer of thallium sulphides is formed on the surface of the sulphur layer. Deposition of thallium sulphide films from the solution of higher polythionic acid on the LDPE films is not homogeneous and has the separated island type morphology.

The inhomogeneous film structure is related to the LDPE film surface structure and islands of different PE crystallization direction. The treatment of LDPE films with the sulphur layer in the Tl₂SO₄ solution does change significantly the structure and roughness of the films, but the average roughness and film thickness are higher when compared with the sulphur films.

**XRD characterization**

The phase composition of thallium sulphides could be qualitatively and semiquantitatively characterized by the X-ray diffraction. It is not easy to identify the structure of thallium sulphide layers on the polymeric surface due to the peculiarities of composition materials: polycrystallinity of layers; existence of several phases of TlₓSᵧ, which have different composition and structure; a crystal structure of a polymeric film. Despite the mentioned difficulties, the phasic composition of the layer of thallium sulphides on the PE surface was analysed by the method of X-ray diffraction analysis [34, 37, 59].

The peak of the monoclinic TIS phase predominates in the diffractograms of obtained TlₓSᵧ layers when the LDPE films were sulphured in the solution of H₂S₃O₆ during 15, 60 and 120 min at 60°C temperature (Fig. 10). A strong peak of the TIS(223) phase is seen in the diffractograms at 2θ = 24.18° (corresponding to interplanar spacing d = 3.69 Å). The peak of the TIS(130) phase at 2θ = 25.6° (corresponding to d = 3.48 Å) is also observed in the diffractograms of these samples. The peaks of another two thallium sulphide phases in the diffractograms of LDPE samples initially sulfured during 120 min are observed (Fig. 10, Curve 3): tetragonal Tl₂S₄ at 2θ = 11.9, 31.2 and 46.9° (corresponding to d = 7.3, 2.87 and 1.955 Å) and rhombohedral Tl₂S at 2θ = 29.4 and 49.36° (corresponding to d = 2.9 and 1.79 Å).

**Fig. 10.** X-ray diffraction patterns of thallium sulphide layers on the surface of LDPE. LDPE initially treated with a solution of H₂S₃O₆ at 60°C at different time and then with the Tl₂SO₄ solution at 80°C for 10 min. The sulphurisation time, min: 15 (1), 60 (2) and 120 (3)
The phasic composition of the \( \text{Tl}_x\text{S}_y \) layers, made by repeating cycles of sulphurisation and treatment with the thallium(I) salt solution for three times, was also identified by the X-ray diffraction analysis. The research results showed that by repeating the treatment cycles of LDPE, three phases of thallium sulphides were identified in the diffractograms: TIS, \( \text{Tl}_x\text{S}_y \) and \( \text{Tl}_2\text{S}_3 \). In the diffractograms the peaks of TIS and \( \text{Tl}_2\text{S}_3 \) phases, getting more intensified when the duration of LDPE sulphurisation is prolonged and the number of treatment cycles is increased, are most intensive.

XPS characterization

Later the coat surface is etched by Ar\(^+\) ions up to 1 nm depth and again it is investigated. This analysis is limited only in the research of the chemical and phasic composition of a very thin (1 nm) \( \text{Tl}_x\text{S}_y \) layer as the thickness of all layer reaches several tens of \( \mu \text{m} \). Considering the data obtained by the XPS method – atomic percentage of elements, bond energies and separate elements \( \text{Tl}_4\text{f}_{7/2}, \text{O} 1\text{s} \) and \( \text{S} 2\text{p} \) spectra – we were seeking to identify the composition of the \( \text{Tl}_x\text{S}_y \) layers [58].

By means of the XPS method we identified that on the polymer surface the layer consisting of different thallium, sulphur and oxygen compounds is formed. The following compounds were identified: TIS, \( \text{Tl}_x\text{S}_y \), \( \text{Tl} (\text{OH})_y \), \( \text{S}_8 \) and \( \text{Tl}_2\text{O}_y \). The study of \( \text{Tl}_x\text{S}_y \) layers revealed that oxygen (30–57 at.%) made the largest part on the surface of all samples of the PE film. During the formation of \( \text{Tl}_x\text{S}_y \) layers on the PE surface all processes take place in the open environment, so it is impossible to avoid environment influence upon the chemical composition of the obtained layers and on the surface of thallium sulphide layers oxygen can always be detected. As metal sulphides, when getting them in the matrix of the polymer surface by the sorption–diffusion method, are formed in the shape of dendrites, so among the dendrites one can detect adsorbed hardly soluble \( \text{Tl}_x\text{SO}_y \). In addition, if the oxidation–reduction reaction when thallium(I) ions react with element sulphur occurs, thallium(III) ions make insoluble \( \text{Tl} (\text{OH})_y \) together with the present groups of \( \text{HO}^- \) in alkali-

ized \( \text{Tl}_x\text{SO}_y \). In deeper layers oxygen can be interlinked into \( \text{Tl}_2\text{O}_y \). The layers of \( \text{Tl}_x\text{S}_y \) on the polymer surface are formed in islets because the diffusion of sulphur and thallium(I) ions is possible only in amorphous areas of PE, and atmospheric oxygen faces no difficulties to approach the latter.

When the surface of the layer is etched by Ar\(^+\) ions, the amount of oxygen considerably decreases (0–20 at.%). This proves that in the mentioned compounds a greater part of oxygen is found only on the surface.

The data of XPS analysis show that the composition of the \( \text{Tl}_x\text{S}_y \) layers formed under different conditions is very similar. Thus the research performed by the RAEF method confirms the results obtained by the X-ray diffraction research of the \( \text{Tl}_x\text{S}_y \) layers – the formation of thallium sulphides of different phases in the layers of \( \text{Tl}_x\text{S}_y \) on the PE surface.

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AUKŠTOSIOS POLITIONATO RŪGŠTYS – PIRMTAKAI SUDARANT TALIO SULFIDŲ ŠLUOKSNIUS POLIETILENO PAVIRŠIUI

Santrauka
Patobulintas ir tyrinėtas talio sulfidų plonų sluoksnii nusodinimo ant žemo tankio polietileno (LDPE), naudojant aukštųjų politionato rūgščių (H3O+), metodas. Įvedus sieringumo sąlygas, sierintas polimeras veikiamas Tl(I) tirpalu, siera, esanti po tirpalo temperatūrą, jos sieringumą ir išlaikymo tirpale trukmę.

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