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MORPHOLOGY, PHASE AND CHEMICAL COMPOSITION OF THE NANOSTRUCTURES FORMED IN THE SYSTEMS CONTAINING LANTHANUM, CERIUM, AND SILVER

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X-ray phase and thermogravimetric analysis, scanning electron microscopy and energy-dispersion spectroscopy were used to study the products of phase formation during the precipitation of lanthanum and cerium salts in the presence of silver nitrate and recipients of precipitators, nucleating agents and hydrolysis regulators. Thermogravimetric analysis shows the completion of the $\text{La}(\text{OH})_3$ lattice dehydroxylation process at a temperature of $\sim 300^\circ\text{C}$ and probable destruction of sulfates at a temperature of $\sim 340^\circ\text{C}$. The phase interaction of lanthanum oxide(III) with silver ends at $T \sim 400^\circ\text{C}$. The DTG curve shows a two-stage weight loss, which characterizes the destruction of lanthanum and silver hydroxides (250°C) and the removal of sulfates ($\sim 340^\circ\text{C}$), respectively. According to the TG, the total weight loss is 21.6%. For the cerium-containing system the only endothermic effect of dehydroxylation of cerium hydroxide at $T = 250^\circ\text{C}$ with its conversion into cerium dioxide is observed. The destruction of nitrates (anionic component of solutions) takes place at the temperature of 400°C . Weight loss takes place at $T = 150^\circ\text{C}$ and is 53.9%. Thus, on the basis of TG-DTA data, it can be assumed that the formation of composites particles based on lanthanum and cerium oxides, modified with silver, ends at the temperature of 400°C . The X-ray diffraction data shows that at the initial stage the system undergoes the formation of cerium and lanthanum hydroxides, and during lyophilization of the precipitate ($T = 160^\circ\text{C}$) the crystal lattice of hydroxides partial dehydroxylation takes place with the formation of trigonal oxides La_2O_3 and Ce_2O_3 . It has been found that the presence of silver cations in the solution can affect the phase composition of lyophilized structures and the formation of the CeO_2 phase. It is shown that the hydroxylamine chloride injection into the system can initiate the silver restoration on the lanthanum oxide surface and also partially restore it to the LaO phase. Temperature treatment of the samples ($T = 400^\circ\text{C}$) promotes homogenization of the precipitate composition: formation of 30 nm cerium dioxide particles with silver clusters evenly distributed on its surface, and hexagonal lanthanum oxide plates with individual silver particles as the second phase. In three-component systems, two modifications of lanthanum oxides (trigonal and cubic), cerium dioxide and metallic silver are formed. It is found that the chemical composition of the precipitates contains the main elements – La, Ce, O, Ag and impurity – S or Cl, as the anionic component of the initial solutions, N and K in the composition of the initial suspension. It is shown that the morphology of the samples is represented by hexagonal structures of lanthanum hydroxide and oxide, spherical and pseudocubic particles of cerium dioxide and lanthanum oxide, spherical clusters of silver.

Keywords: trigonal lanthanum oxide, cerium dioxide, doping of REE oxides with silver, phase formation, morphology of cerium and lanthanum oxides, silver

INTRODUCTION

When conducting modern medical and biological research, among the whole variety of oxides of rare earth elements (REE), systems based on cerium dioxide are most often used, for which the special term “nanoceria” has been introduced in the scientific literature [1]. Structure-sensitive properties and biomedical applications of nanodispersed cerium dioxide are shown in paper [2]. Interest in these systems is based on the capability of cerium oxide particles to absorb oxygen and reversible transition between Ce^{3+} and Ce^{4+} cations on their surface,

which is used in catalytic and photocatalytic processes [3, 4]. The use of cerium oxide nanoparticles in biology and medicine is associated, for example, with their auto-catalytic anti-oxidant behavior [5, 6]. Nanoceria is known as free radical scavenger or antioxidant [7] that is distinguished by biocompatibility and provides a significant neuroprotective and regenerative effect that has been studied on rats [8]. Ceria nanoparticles possess unique oxidase-like activity, as they can facilitate the fast oxidation of organic dyes and small molecules under slightly acidic conditions without the need of hydrogen peroxide [9]. The key protective role

of nanoparticles is closely connected with their possibility to absorb harmful UV radiation without scattering the useful visible one and reducing oxidative stress damage, such as that induced by H_2O_2 , thanks to the surface redox Ce^{3+} - Ce^{4+} reactivity of nanoparticles [10]. In biological research, particles of lanthanum oxide are much less widespread, which, nevertheless, exhibit hemocompatibility and antibacterial properties [11]. Lanthanum oxide is also used in the production of fluorescent materials based on hydroxyapatite, which use rare earth elements, including lanthanum (La^{3+}), that have been developed in the biomedical field [12] and as a tracer for the study of tight junctions in epithelia by transmission electron microscopy [13]. Numerous experimental and review works have been devoted to the biomedical application of silver, in particular, [14, 15], however, until now, the question of the mechanisms of its interaction in biological systems including human body [16], toxicity of nanosized silver particles [17], its bactericidal, cytotoxic [18] and therapeutic [16] action remains controversial.

Nanoscale composites based on rare earth oxides modified with various noble metals, including silver, are considered as attractive materials for technical and biomedical applications due to the combination and improvement of their individual intrinsic properties. In particular, the decoration of cerium dioxide particles with silver makes it possible to obtain materials with improved photocatalytic properties as a charge carrier and the improvement of photocatalysis performance. The decoration of Ag nanoparticles on CeO_2 nanorods can also prevent the recombination of e⁻-h⁺ pairs by efficient electron transfer from CeO_2 to Ag [19]. Structures of composite particles Ag – La_2O_3 in which silver particles are localized on the surface of lanthanum oxides, and are restored from solution using sodium borohydride as a reducing agent, showed anti-oxidizing and significant antibacterial effect *in vitro*. Thus, the obtained composites can be potentially extended to develop drug-labels and other antibacterial agents [20, 21]. Silver-cerium nanoparticles, synthesized using the sol-gel method from solutions of cerium nitrate and silver, which are spherical silver particles coated with nano cerium showed higher inhibitory properties compared to uncoated bare silver particles. The disinfection effect of Ag-doped

CeO NP was more pronounced on *Staphylococcus aureus* than *Escherichia coli*, although the difference was not wide. The catalytic effect of cerium oxide particles obtained using an inexpensive precipitation method and evaluated their disinfecting aptitudes with the turbidimetric and plate count methods was used for cytotoxicity toward microorganisms at room temperature [22]. Wherein the silver-doped ceria exhibits the highest sterilization capability, yet the undoped ceria is the most intriguing. The disinfection effect of undoped ceria is moderate in magnitude, demanding a physical contact between the ceria surface and bacteria cell wall, or the redox catalysis that can damage the cell wall and result in the cell killing. Evidently, this effect is short-range and depends strongly on dispersion of the nanoparticles. In contrast, the disinfection effects of silver-doped ceria reach out several millimeters since it releases silver ions to poison the surrounding microorganisms. It should be noted that particles of this type obtained by a facile oxidation-reduction reaction method have been characterized as electro-catalysts for the air cathode of lithium-air batteries operated in a simulated air environment. [23]. The photocatalytic properties of the novel Ag decorated by CeO_2 catalyst that are effective under sunlight upon the textile pollutant *Rose Bengal* destruction are considered in the work [24]. It was found that great enhancement of the degradation efficiency for Ag/ CeO_2 compared to pure CeO_2 can be ascribed mainly due to decrease in its band gap and charge carrier recombination rate. The possible degradation mechanism has been proposed, it shows the effect of generation of oxygen vacancies owing to the decoration of Ag on the CeO_2 surface. The role of silver clusters in enhancing the photocatalytic activity of cerium oxide particles obtained by the hydrothermal method is indicated in the article [25]. As-prepared Ag NC/ CeO_2 NPs have been employed as photocatalyst to study degradation of dye *Acridine red* under UV light illumination. Results showed that photocatalytic activity of the Ag NC/ CeO_2 was enhanced several times compared to pure CeO_2 NP even with very small amount loading of Ag NC (~3%). The mechanistic interpretation of the increased catalytic activity of Ag NC/ CeO_2 NPs was nicely corroborated by involving participation of

reactive oxygen species. Thus, Ag NC opens a convenient path for improving photocatalytic activity of CeO₂ NP. Catalysts of this kind are used in the carbon oxidation process [26].

Particles of composites based on lanthanum oxide doped with cerium dioxide also exhibit catalytic properties [27]. Both citrate and solvothermal methods allowed the incorporation of Ce into the La₂O₃ lattice as substitution ions on La creating interstitial oxygen species leading to an abundance of electrophilic oxygen species on the catalyst surface. The citrate method originated O⁻ and O₂²⁻ oxygen species, while the solvothermal method produced O⁻ oxygen species. These results can be related to the different types of cerium ions (Ce³⁺ or Ce⁴⁺) on the oxide surface. The catalyst prepared by the solvothermal method showed higher concentration of [Ce³⁺]. The O⁻ oxygen species predominant in the La_{0.75}Ce_{0.25}O (SM) catalyst would be responsible for higher C₂ yield (by 10.5 %) at 750 °C in the oxidative coupling of methane. Generally, it is well known that the synthesis procedure strongly effects on the structure, phase and chemical composition, morphology and physicochemical properties of nanomaterial chemical composition. The aim of the present work is studying the influence of the composition of the precursor species and synthesis procedure on the phase and chemical composition of nanostructures formed in lanthanum- and cerium-based systems doped with silver. In this case the nucleation mediate stage and primary (precursor) phase growing can be indicated as the main aspect to form homogeneous materials with controlled physical-chemical properties.

OBJECTS AND METHODS OF THE RESEARCH

Particle synthesis was performed by the method of co-precipitation in an alkaline medium from solutions of LaCl₃, La(NO₃)₃, La₂(SO₄)₃, Ce(NO₃)₃, and AgNO₃ when adding substances of nucleating agents and hydrolysis regulators to the system. To obtain particles of nanocomposites based on lanthanum and cerium oxides with dopant substances it was selected Ag⁺ (for both oxides) and Ce³⁺ (for lanthanum oxide) in the mass ratio of 2 and 4 wt. %, due to the recommended optimum of 3 wt. % for structures designed for photocatalytic reactions [25]. The precipitates were dried in an oven at

$T = 160$ °C, and then calcined at $T = 400$ °C for 5 h. In addition, silver was restored from AgNO₃ solution to the surface of pre-synthesized particles of cerium and lanthanum oxides, and the composition and morphology of three-component composite structures based on lanthanum oxide modified with cerium and silver were obtained and studied at a mass content of 2 and 4 wt. %.

The samples were visualized by scanning electron microscopy (SEM). Electron micrographs of the samples were recorded on a MIRA3 TESCAN scanning electron microscope. Determination of the phase composition of the obtained samples was performed by the method of powder X-ray diffraction phase analysis on the DRON-3 device with radiation of the copper anode (CuK_α). The scanning step was 0.05–0.1 degrees, exposure – 4 s, range of 2θ angles – from 15 to 90°. Samples were taken at standard temperature. The International Powder Standards Committee (JSPDS International Center for Diffraction Data 1999) database was used to determine the phase composition. A simultaneous study of thermogravimetric and differential thermal properties (TG-DTA) of the composite particles were performed in the static air atmosphere by a derivatograph Q-1500D (Hungary). The parameters of the pattern recording were as follows: the samples 200 mg were heated at the rate 10 °C/min from 20 to 1000 °C; the sensitivity was 20 mg; TG – 500, DTG – 500, and DTA – 250. The samples were placed into a corundum crucible and covered by a quartz beaker to create a homogenous temperature field.

RESULTS AND DISCUSSION

Characteristics of the primary mineral phases formed at $T = 160$ °C. The phase composition study of lyophilized products of co-precipitation of lanthanum and cerium salts with silver nitrate at a mass content of c(Ag⁺) of two and four wt. percent indicates the mainly hydroxide structures formation in the sediment, respectively, La(OH)₃ (# 13-0084) (Fig. 1 a) and Ce(OH)₃ (# 19-0284) (Fig. 1 c). Note that the diffractogram of the lanthanum-containing sample contains reflexes of two phases of lanthanum oxides: La₂O₃ (# 40-1281 / 02-0688) and LaO (# 33-0716) and silver nitrate (# 6-0363). The lyophilized precipitate formed in the cerium-

containing system also contains oxides Ce_2O_3 (# 44-1086) and CeO_2 (# 44-1001), as well as silver nitrate. For a system of lanthanum-containing system containing 2 and 4 wt. % of

cerium in addition to the phase of hydroxide (# 36-1481) and oxide (# 22-641) of lanthanum, CeO_2 reflexes (# 34-0394) are observed.

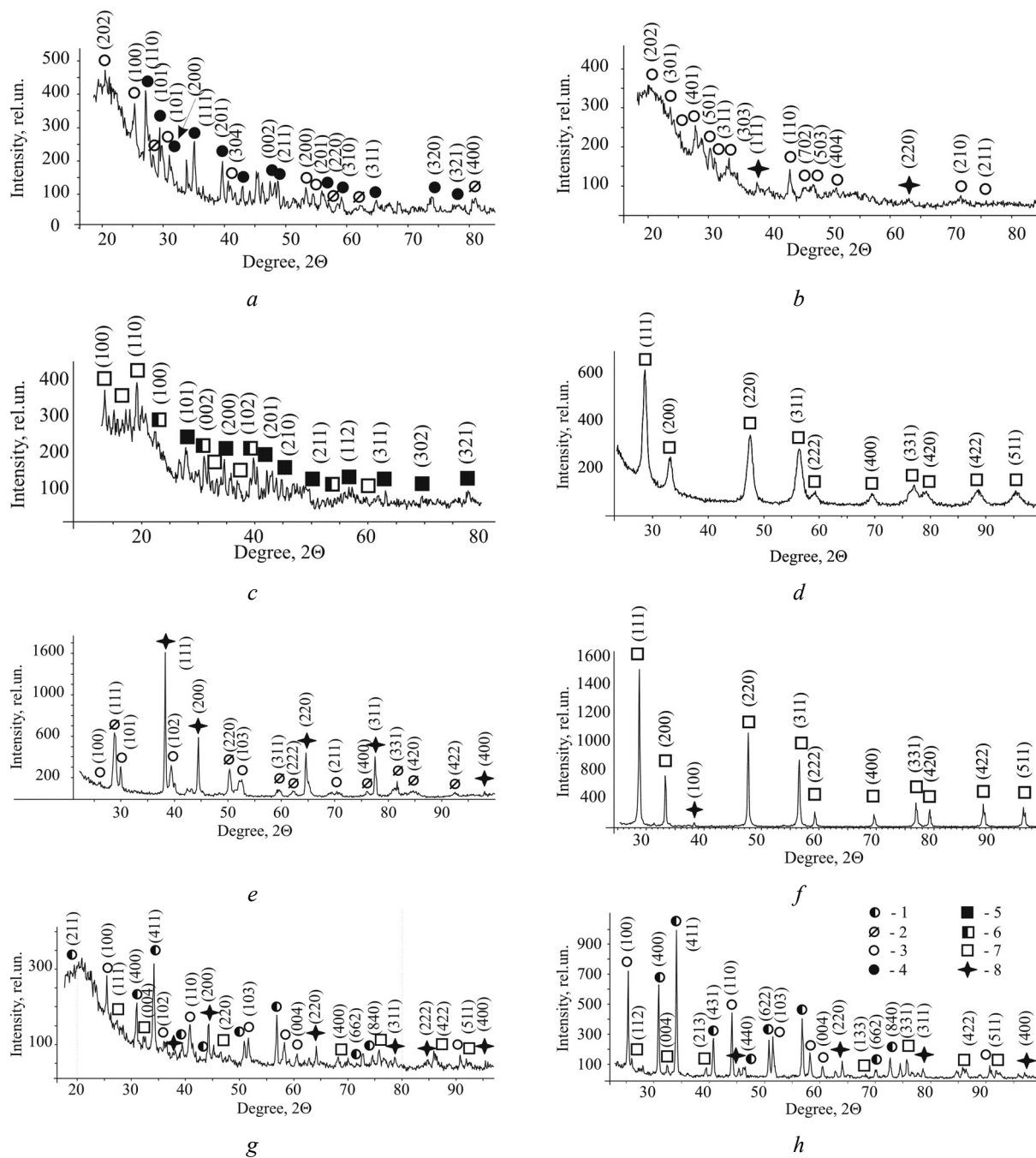


Fig. 1. XRD patterns of the samples formed in the systems: *a* – $\text{La}_2(\text{SO}_4)_3\text{-AgNO}_3$ (160 °C), *b* – $\text{La}_2(\text{SO}_4)_3\text{-AgNO}_3$ (400 °C, 5 h); *c* – $\text{Ce}(\text{NO}_3)_3\text{-AgNO}_3$ (160 °C), *d* – $\text{Ce}(\text{NO}_3)_3\text{-AgNO}_3$ (400 °C, 5 h); *e* – $\text{La}(\text{NO}_3)_3\text{-Ce}(\text{NO}_3)_3$ (160 °C); *f* – $\text{La}(\text{NO}_3)_3\text{-Ce}(\text{NO}_3)_3$ (400 °C, 5 h); *g* – $\text{La}_2\text{O}_3\text{-CeO}_2\text{-Ag}$ 2 wt. % (400 °C, 5 h); *h* – $\text{La}_2\text{O}_3\text{-CeO}_2\text{-Ag}$ 4 wt. % (400 °C, 5 h). Numbers correspond to phases: 1 – La_2O_3 (P3m1); 2 – La(OH) (F); 3 – La_2O_3 (Ia3); 4 – $\text{La}(\text{OH})_3$ (P63/m); 5 – $\text{Ce}(\text{OH})_3$ (P63/m); 6 – Ce_2O_3 (P321); 7 – CeO_2 (Fm3m); 8 – Ag^0

Fig. 2 shows the SEM images of structures formed as part of lyophilized lanthanum and cerium-containing sediments. Whereas the hexagonal crystals of $\text{La}(\text{OH})_3$ belongs to $P6_3m$ space group (Fig. 2 *a*) and the plate like 2D crystals of $\text{Ce}(\text{OH})_3$ (Fig. 2 *b*) are formed in the presence of 4 wt. % AgNO_3 , an addition of $\text{Ce}(\text{NO}_3)_3$ leads to change in the morphology of lanthanum hydroxide (Fig. 2 *c*). The average chemical composition of $\text{La}(\text{OH})_3$ according to

EDS-data includes wt. %: 63 – La, 24 – O, and 13 total N, S, K, but Ag is undetectable. The EDS spectrum of $\text{Ce}(\text{OH})_3$ shows, wt. %: 66 – Ce, 3.8 – Ag, 23 – O, and 7.2 – N. The composition of $\text{La}(\text{OH})_3$ formed in the presence of Ce^{3+} is as follows, wt. %: 47 – La, 5 – Ce, 35 – O, and 13 – total N, Na. Therefore, the nature of rare earth species and doping elements determines the morphology and composition of primary phase.

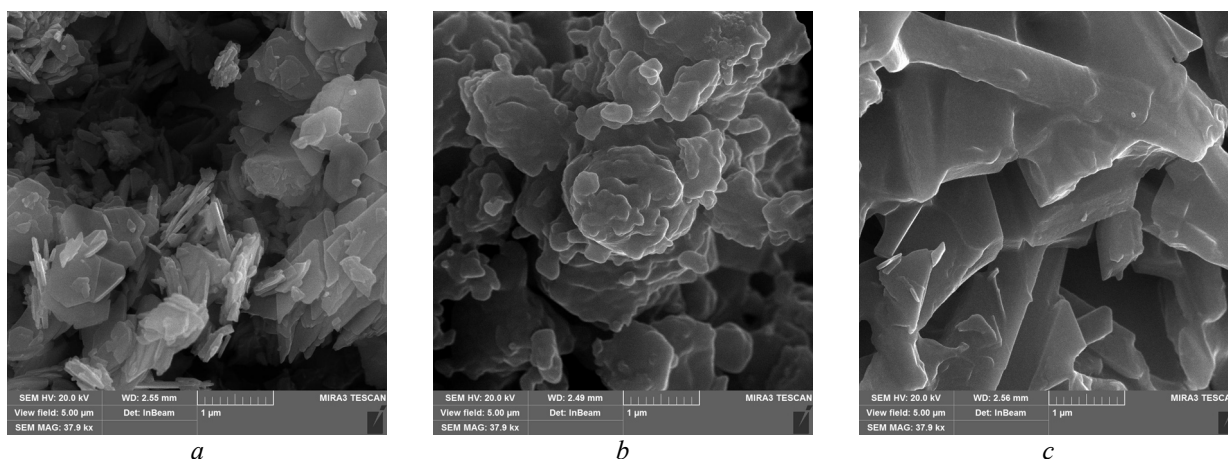


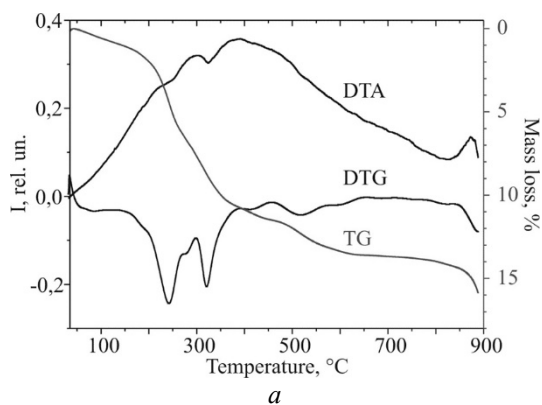
Fig. 2. SEM images of the precursor phases: *a* – $\text{La}(\text{OH})_3$ doped with Ag^+ ; *b* – $\text{Ce}(\text{OH})_3$ doped with Ag^+ ; *c* – $\text{La}(\text{OH})_3$ formed in the presence of Ce^{3+}

Fig. 3 shows the thermogravimetric analysis data of the initial samples obtained by co-precipitation of lanthanum and cerium-containing solutions with a solution of silver nitrate with silver concentration of 4 wt. %. There are two endothermic effects on the DTA curve of the lanthanum-containing sample (Fig. 3 *a*), that indicate the completion of the $\text{La}(\text{OH})_3$ lattice dehydroxylation process at a temperature of $\sim 300^\circ\text{C}$ and probable destruction of sulfates at a temperature of $\sim 340^\circ\text{C}$. The phase interaction of lanthanum oxide(III) with silver ends at $T \sim 400^\circ\text{C}$. The endothermic peak at $T = 900^\circ\text{C}$ can be attributed to the change in the crystalline modification of lanthanum oxide. The DTG curve shows a two-stage weight loss, which characterizes the destruction of lanthanum and silver hydroxides (250°C) and the removal of sulfates ($\sim 340^\circ\text{C}$), respectively. Destruction of nitrate is observed at $T = 380^\circ\text{C}$. According to the TG, the total weight loss is 21.6 %. Taking into account that during the preliminary preparation the samples were subjected to heat treatment at $T = 160^\circ\text{C}$,

there is no reflex on the DTG curve, which characterizes the loss of adsorption-bound water. For the cerium-containing system (Fig. 3 *b*) the only endothermic effect is observed of dehydroxylation of cerium hydroxide at $T = 250^\circ\text{C}$ with its conversion into cerium dioxide. The destruction of nitrates (anionic component of solutions) takes place at the temperature of 400°C . Weight loss takes place at $T = 150^\circ\text{C}$ and is 53.9 %. Thus, on the basis of TG-DTA data, it can be assumed that the formation of composite particles based on lanthanum and cerium oxides, modified with silver, ends at the temperature of 400°C .

Characteristics of the products of phase transformation at $T = 400^\circ\text{C}$. Heat treatment of lyophilized samples at the temperature of 400°C leads to a change in their phase composition (Fig. 1 *b*). In particular, the lanthanum-containing system has a dominant phase of La_2O_3 oxide (# 05-0602). The diffraction pattern shows two weak reflexes of metallic silver (# 4-0783). In the cerium-containing system (Fig. 1 *d*), the formation of homogeneous

nanosized particles of cerium dioxide (# 34-0394) takes place. Dehydrated samples calcination at 400 °C for 5 h leads to the change in the particle morphology. In particular, hexagonal particles of lanthanum hydroxide are transformed into a trigonal (rhombohedral) structure of lanthanum oxide (silver clusters are restoring on its surface)



(Fig. 4 a). The formation of cerium dioxide aggregates with evenly distributed small clusters of silver (Fig. 4 b) is observed in the cerium-containing system. In the lanthanum-containing system, the presence of cerium (4 wt. %) is seen in the nanosized CeO₂ particles formed on the edges and planes of La₂O₃ plates (Fig. 4 c).

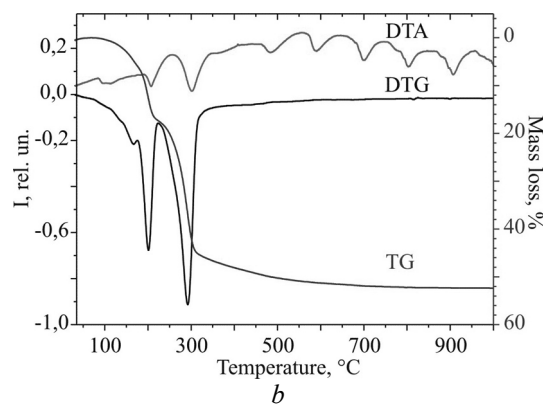


Fig. 3. TG-DTA curves of the primary samples formed via precipitation of RE salt solutions with AgNO₃ in the light alkaline media in the systems: *a* – La₂(SO₄)₃-Ag⁺; *b* – Ce(NO₃)₃-Ag⁺

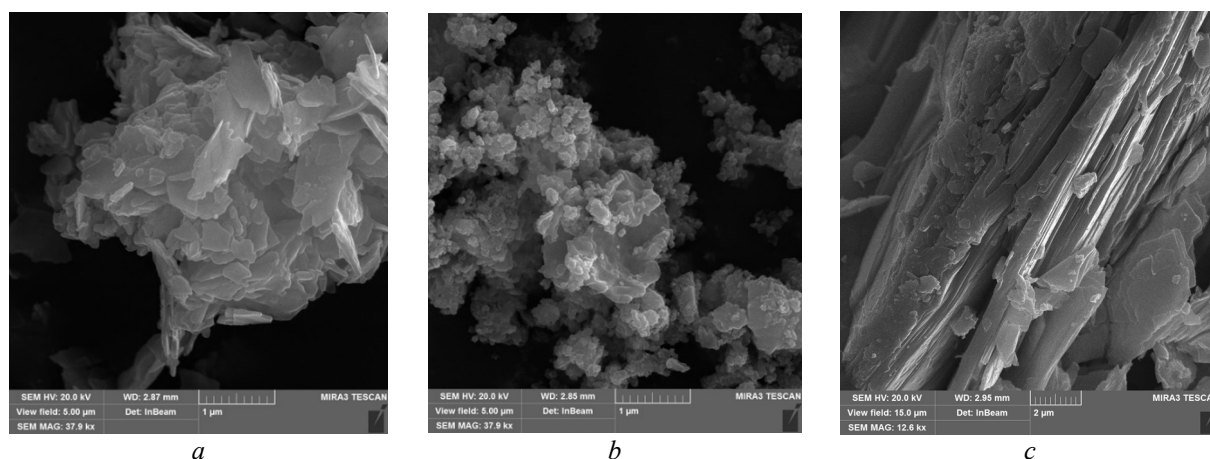


Fig. 4. SEM images of structures formed in systems at $T=400$ °C: *a* – lanthanum sulfate - silver nitrate; *b* – cerium nitrate- silver nitrate; *c* – lanthanum sulfate - cerium nitrate

According to EDS spectroscopy, for the procedure of synthesis the use of lanthanum sulfate leads to the insertion of sulfur into the sample. For the system *lanthanum sulphate - silver nitrate* 2 wt.%, the average composition of the sample is: O – 35, S – 11.8, K – 0.6, Ag – 1.3, La – 51.3. For the system *cerium nitrate - silver nitrate* 4 wt.% the composition of the sample wt. % is: O – 13.6, Ag – 8.5, Ce – 77.9. In the system of *lanthanum nitrate - cerium nitrate* with the content of the last 4 wt. % a precipitate is formed, the chemical composition

of which contains wt. %: 56 – La; 3.4 – Ce; 28 – O; 3 – Na; 9.6 – N.

Characteristics of the products of red-ox process on oxide surfaces and next sintering of the samples at $T=400$ °C. Particles of previously formed oxides lanthanum and cerium ($T=400$ °C) were used to model the process of silver reduction from AgNO₃ solution. A solution of hydroxyl amine chloride was chosen as the reducing agent. The obtained samples were subjected to additional heat treatment. According to X-ray diffraction, in the

lanthanum-containing system there are phases of two oxides - trigonal La_2O_3 (# 05-0602) and cubic LaO (F), and a phase of metallic silver (# 6-0363) (Fig. 1 e). For the cerium-containing system, the phase of cerium dioxide (# 34-0394) and metallic silver (Fig. 1 f) was determined. According to EDS data, in the lanthanum-containing system there are two types of structures that differ in their composition: the first type belongs to lanthanum oxides and in its chemical composition contains, wt. % - 72 - La; 10 - Ag; 7.6 - S; 10.4 - O, and the second - reduced silver, wt. %: 96.6 - Ag; 3.4 - O. For a

cerium-containing system is seen a uniform distribution of chemical elements in the composition of the sample, which is, wt. %: 81 - Ce; 3 - Ag; 16 - O. The SEM images show the morphology of rhombohedral lanthanum oxide particles (Fig. 5 a) and spherical silver particles (clusters) on the surface of La_2O_3 plates (Fig. 5 b). The morphology of the structures of the cerium-containing system is shown in (Fig. 5 c). It is a spherical particle that forms aggregates and is localized on compacted aggregates of much larger size.

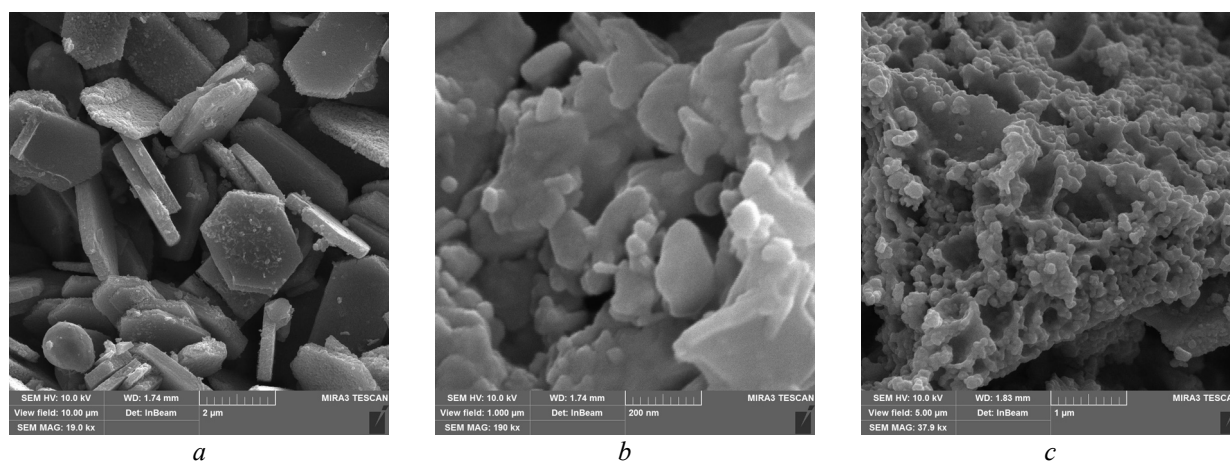


Fig. 5. Morphology of structures formed during the silver restoration from AgNO_3 solution to the surface of oxide particles: *a* – trigonal lanthanum oxide; *b* – spherical particles (clusters) of silver on the surface of lanthanum oxide; *c* – spherical particles and aggregates of cerium dioxide with silver clusters on the surface

Characteristics of phase formation products in three-component $\text{La}_2\text{O}_3\text{-CeO}_2\text{-Ag}$ systems.

Consideration of the phase composition of powders obtained in three-component systems based on lanthanum oxide, doped with cerium and silver, indicates that in systems with dopant' 2 and 4 wt. % there are two lanthanum oxides belonging to the cubic syngony, spatial group Ia3 and trigonal syngony with the spatial group P3m1 (JCPDS file No. 02-0688), cerium dioxide with the spatial group Fm3m (JCPDS file No. 34-0394) and reduced silver (JCPDS file No. 04-0783). The difference is in the ordering (crystallinity degree) increasing for the sample with 4 wt. % of dosing impurities (Fig. 1 h), compared with the sample, which contains 2 wt. % of cerium and silver (Fig. 1 g).

At the same time, according to EDS data, the chemical composition of the samples contains, wt. %: 62 - La; 4 - Ag; 10 - O; 24 - Cl, while the phase of cerium dioxide is not identified. The

use of lanthanum chloride for the sample synthesis leads to the chlorine presence in the sediment composition.

The morphology of the structures formed in the lanthanum oxide system, modified with cerium and silver in the ratio of 2 and 4 wt. %, is shown in Fig. 6. In Fig. 6 a the image is seen of deformed plates of lanthanum oxide, on the surface of which there are spherical nanometric particles of cerium dioxide and clusters of restored silver. In Fig. 6 b the enlarged image of such structures can be seen, and the difference in the sizes of the components of the three-phase dispersion is well observed. Clusters of silver on the lanthanum oxide surface are shown in Fig. 6 c.

Thus, a study of morphology, phase and chemical composition of nano and micro-dimensional structures formed in the systems of inorganic salts of lanthanum, cerium nitrate and argentum, allowed us to determine the specifics

of the phase formation process dependent on the chemical composition of the initial solutions, heat treatment of sediments and adding dopants, in particular, silver. Analysis of the cerium-containing system shows that at the initial stage of synthesis, which consists in the formation and lyophilization of the precipitate at $T = 160\text{ }^{\circ}\text{C}$, the composition of the precipitate is mainly

cerium hydroxide(III). At the same time, the input of potential reducing agents into the suspension – the admixture of silver nitrate and sodium hydroxide, promotes the spontaneous course of the redox process – oxidation of Ce^{3+} to Ce^{4+} , with the formation of CeO_2 particles and nanoparticles or Ag^0 clusters that was shown in [23].

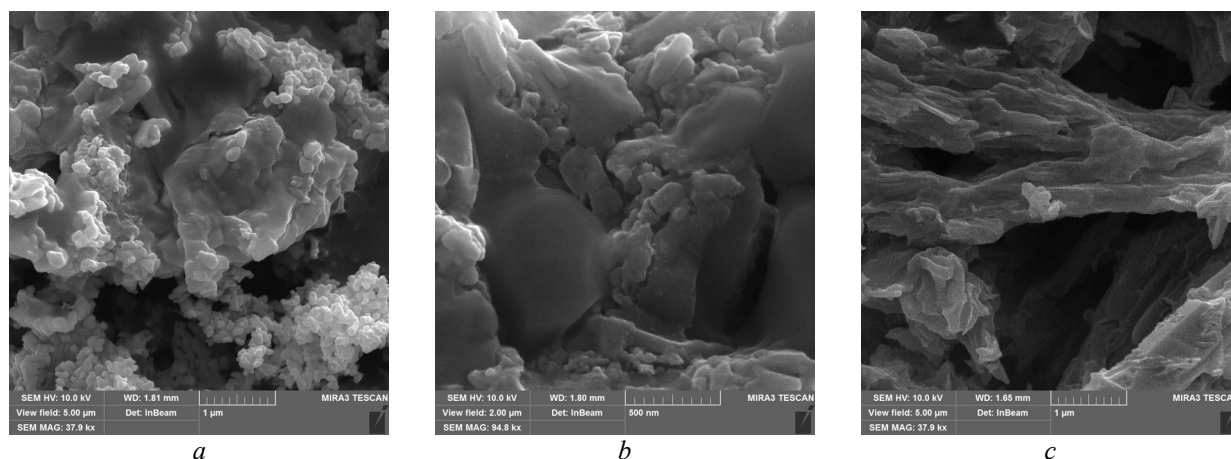


Fig. 6. Morphology of structures formed in the three-component $\text{La}_2\text{O}_3\text{-CeO}_2\text{-Ag}$ system: *a* – the sample morphology general view; *b* – deformed plates of lanthanum oxide with spherical nanoparticles of cerium dioxide; *c* – clusters of silver on the surface of lanthanum oxide

Drying of the jelly-like precipitate at $T = 160\text{ }^{\circ}\text{C}$ promotes the beginning of the $\text{Ce}(\text{OH})_3$ crystal lattice dehydroxylation with the formation of a slight impurity of the Ce_2O_3 phase. Subsequent heat treatment of the lyophilized sample leads to homogenization of the polyphase precipitate with the formation of spherical CeO_2 particles, size $\sim 30\text{ nm}$, with evenly distributed on its surface clusters of silver. Silver restoration from AgNO_3 solution in the presence of hydroxyl amine with subsequent calcination of the sample for 5 h at $T = 400\text{ }^{\circ}\text{C}$ leads to particle size increase of cerium dioxide and the appearance of a metallic silver reflex on the diffraction pattern. According to EDS, such a precipitate contains only such components as Ce, Ag, O.

The lyophilized precipitate of the lanthanum-containing system consists mostly of lanthanum(III) hydroxide structures, at the same time, the same as in the cerium-containing system, partial dehydroxylation of the $\text{La}(\text{OH})_3$ crystalline structure to the La_2O_3 formation and the presence of LaO impurity is observed. This is clearly seen by the reflexes on the diffraction

pattern (Fig. 1 *c*). In the initial samples of both systems, the presence of silver-containing compounds is not identified due to their low content (2 and 4 wt. %) and the sensitivity limit of X-ray diffraction being equal to 5 % by weight of the sample. Heat treatment of the sample ($400\text{ }^{\circ}\text{C}$) leads to its homogenization with formation of the dominant phase of La_2O_3 in trigonal syngony with spatial group $P3m1$. When a solution of silver nitrate and a reducing agent is inputted into the lanthanum oxide La_2O_3 (Ia_3) system, a second phase of lanthanum oxide LaO (F) appears on the sediment, probably localized on the surface of trigonal La_2O_3 oxide particles, and silver clusters. In the chemical composition, according to the EDSF data, in the original sample, in addition to lanthanum and oxygen, impurities N, S, K are identified, but the presence of Ag is not detected. After the sample calcination at $T = 400\text{ }^{\circ}\text{C}$, La, O, Ag and S remain in it. The same elemental composition is saved in the sample of lanthanum oxide with restored silver, but in this case the presence of lanthanum and silver-containing phases is clearly traced.

The study of three-component systems in which the lanthanum-containing phase was supplemented by a cerium and silver in a mass dopants ratio 2 and 4 wt.% indicates the presence of trigonal and cubic lanthanum oxides, a small content of cerium dioxide and metallic silver. The elemental composition is stable and contains wt. %: La, Ce, O, Cl. The presence of chlorine is explained by the use of LaCl_3 for sample synthesis. It is obvious that the anionic component of the dispersion medium is a part of the precipitate structure, so the choice of the anionic component of the salt is an important aspect in the further usage of such structures. In particular, chloride in contrast to sulfate and nitrate is the part of physiological solution (0.9% NaCl) it seems promising to apply chloride solutions for the materials synthesis for medical and biological purposes.

CONCLUSION

The study of phase formation processes in the systems of inorganic cerium and lanthanum salts in the presence of silver nitrate and excipients – precipitators, nuclei and hydrolysis regulators shows that at the initial sludge drying stage the $\text{Ce}(\text{OH})_3$ or $\text{La}(\text{OH})_3$ phases dominate in its composition. Lyophilization of the precipitate at the temperature of 160 °C leads to partial dehydroxylation of the crystal lattice of hydroxides with the formation of Ce_2O_3 oxide or trigonal La_2O_3 . At the same time, under the presence of silver and sodium hydroxide cations

in the system, the Ag^0 reduction and Ce^{3+} oxidation with the formation of the CeO_2 phase are probable in the cerium-containing system. At the same time, the lanthanum-containing system is characterized by the formation of an additional phase of LaO , it is most evident when one adds a reducing agent – hydroxyl amine chloride to the system. Calcination of the samples at $T = 400$ °C leads to homogenization of the precipitate composition – formation of 30 nm particles of cerium dioxide with evenly distributed on their surface clusters of silver, and lanthanum oxide with individual reduced silver nanoparticles. When silver is reduced on the surface of previously obtained oxides of cerium and cubic lanthanum, the particle size of CeO_2 increases, and lanthanum is partially reduced to the LaO particles formation. The formation of two modifications of lanthanum oxides (cubic and trigonal), cerium dioxide and metallic silver is obvious for three-component systems based on lanthanum oxide doped with cerium and silver. From the morphology point of view, hexagonal plates of lanthanum hydroxide and oxide, spherical and close to the cubic shape of lanthanum oxide and cerium dioxide particles, as well as spherical clusters of silver are formed in the system. Talking about the chemical composition, in addition to the main chemical elements Ce, La, O, Ag, the systems have S or Cl, dependent on the anionic component of the lanthanum salt, as well as N and K.

Морфологія, фазовий і хімічний склад наноструктур, утворених в системах, які містять лантан, церій і срібло

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Методами рентгенофазового і термогравіметричного аналізу, скануючої електронної мікроскопії та енерго-дисперсійної спектроскопії проведено дослідження продуктів фазоутворення при осадженні солей лантану і церію в присутності нітрату срібла і допоміжних речовин осадників, зародкоутворювачів і регуляторів гідролізу. Термогравіметричний аналіз свідчить про те, що процес дегідроксилювання кристалічної ґратки $\text{La}(\text{OH})_3$ закінчується за температури ~ 300 °C, а вірогідна деструкція сульфатів відбувається за температури ~ 340 °C. Фазова взаємодія оксиду лантану(III) з сріблом закінчується за $T \sim 400$ °C. На кривій ДТГ спостерігається два рефлекси втрати маси, які характеризують руйнування структури гідроксидів лантану та срібла (250 °C) та видалення сульфатів (~ 340 °C), відповідно. Згідно з даними ТГ, сумарна втрата маси становить 21.6 %. Для церієвмісної системи простежується єдиний

ендотермічний ефект дегідроксилювання гідроксиду церію за $T = 250\text{ }^{\circ}\text{C}$ та його перетворення на фазу діоксиду церію. Руйнування нітратів (аніонна складова розчину) відбувається за температури $400\text{ }^{\circ}\text{C}$. Втрата маси простежується за $T = 150\text{ }^{\circ}\text{C}$ та становить 53.9 %. Таким чином, на підставі даних ТГ-ДТА встановлено, що утворення частинок композитів на основі оксидів лантану і церію, модифікованих сріблом, закінчується за температури $400\text{ }^{\circ}\text{C}$. Згідно даних РФА, на вихідному етапі в системі триває формування гідроксидів церію і лантану, а при ліофілізації осаду ($T = 160\text{ }^{\circ}\text{C}$) часткове дегідроксилювання кристалічної ґратки гідроксидів з утворенням оксидів тригонального La_2O_3 і Ce_2O_3 . Встановлено, що наявність в розчині катіонів срібла може впливати на фазовий склад ліофілізованих структур і сприяти утворенню фази CeO_2 . Показано, що введення в систему хлориду гідроксиламіну може не тільки ініціювати відновлення срібла на поверхні оксиду лантану, але також частково відновлювати його до фази LaO . Температурна обробка зразків ($T = 400\text{ }^{\circ}\text{C}$) сприяє гомогенізації складу осадів: формування 30 нм частинок діоксиду церію із рівномірно розподіленими на його поверхні кластерами срібла, та лусочока тригонального оксиду лантану з наночастинками срібла як другої фази. В трикомпонентних системах утворюються дві модифікації оксидів лантану (тригонального і кубічного), діоксид церію і металічне срібло. Встановлено, що в осадах наявні головні елементи – La , Ce , O , Ag і домішні – S або Cl , як аніонна складова вихідних розчинів. До складу вихідної суспензії входять також слідові кількості N і K . Показано, що морфологія зразків представлена гексагональними структурами гідроксиду лантану і тригональними – його оксиду, сферичними та псевдокубічними частинками діоксиду церію і оксиду лантану, сферичними кластерами срібла.

Ключові слова: тригональний оксид лантану, діоксид церію, допущання сріблом оксидів РЗЕ, фазоутворення, морфологія оксидів церію і лантану, срібло

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