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Formation of polyfunctional photocatalytically active layered oxide materials using coordination nitrates REE and alkali metal as precursors

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The information on alkaline coordination nitrates of rare earth elements of the cerium subgroup - precursors of promising modern multifunctional materials - on the conditions of their formation and existence, the nature of chemical bonding, composition, structure, shape of Ln coordination polyhedra, type of ligand coordination, existence of isotype series is generalized. on stoichiometry of structure, structure, the found out characteristic properties. The obtained data are the basis for detection, identification, control of the phase state of processing objects in the preparatory stages in the formation of nanostructured oxide composite systems of lanthanides and transition elements of special purpose, with catalytic and photocatalytic activity, self-cleaning coating with hydrophilic properties; development of various combined methods of their activation

Keywords: alkaline coordination nitrates of lanthanides, formation conditions, crystal structure of compounds, transformations with physical activation.

Формування поліфункціональних фотокаталітично активних шаруватих оксидних матеріалів з використанням координаційних нітратів РЗЕ та лужних металів як прекурсорів

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Комплексним системним дослідженням взаємодії структурних компонентів у системах нітратів рідкісноземельних і ІА групи елементів періодичної системи – прекурсорів сучасних багатокомпонентних оксидних поліфункціональних матеріалів на їх основі - встановлено утворення чисельного класу лужних координаційних нітратів лантаноїдів. Виявлені об'єктивні закономірності мають фундаментальну і прикладну цінність, поглиблюють розуміння про: хімічні і фізичні властивості Ln, їх комплесоутворюючу здатність; можливість утворення й існування в аналогічних системах асоційованих нових фаз, їх атомно-кристалічну будову, стійкість і стабільність; вплив природи лантаноїдів і лужних металів на структуру комплексних аніонів і сполук у цілому; індивідуальність Ln комплексів; існування ізотипних за складом і структурою груп сполук за природними рядами лантаноїдів і лужних металів; роль NO₃- груп в стереохімії цього класу нітратів; роль води у формуванні найближчого оточення іонів Ln3+- комплексоутворювачів. Одержана система знань про перетворення у системах РЗЕ-вмісних нітратних попередників і кристалохімічні властивості зразків координаційних нітратів Ln набуває особливу цінність при формуванні наноструктурованих шаруватих перовскитоподібних сполук лантаноїдів і перехідних елементів (в тому числі титану та інших), твердих розчинів на їх основі; при встановленні технологічно-функціональних залежностей між способом приготування, варіативністю методу активації технологічних систем, методологією виготовлення цільового продукту і його фазовим складом, параметрами решіток, величиною питомої поверхні, морфологією складових частинок, активністю шарів самоочищуючихся композицій з фотокаталітичними і гідрофільними властивостями, сформованими на поверхні конструкційних і спеціального призначення елементах; при практичній реалізації інноваційних проектів розкладання води для цілей отримання водню, розкладання токсичних органічних речовин у розчинах і повітрі, неповного окиснення вуглеводів; при отриманні нових перовскитоподібних фаз та в інших сферах

Ключові слова: лужні координаційні нітрати лантаноїдів, умови утворення, кристалічна будова сполук, перетворення з фізичною активацією.



Introduction

Currently, the search for methods and complex technologies to create new and improve existing regulations for the production of perfect multifunctional oxide materials of transition and rare earth elements with the structure of defective perovskite, garnet with reproducible properties by low-temperature methods of "soft chemistry" and precursors. They have a complex structure and in scientific and technological terms are difficult objects that are intensively studied [1–11]. Therefore, modern materials science, which is based on them, requires regulatory solutions simple in configuration, low-speed, energy efficient, characterized by scale, and with the ability to reproduce products with a given homogeneity, stability, and a set of predefined characteristics.

Review of the research sources and publications

There are many methods for the synthesis of these oxide multicomponent materials [2–13], based on different physical and chemical principles. The main ones are:

- high-temperature method of solid-phase chemical reactions;
- condensation liquid-phase methods for obtaining nanosized oxide materials based on:
- various variants of mixing of initial components (chemical precipitation (coprecipitation); sol gel; hydrothermal; complexonate homogenization; solvent replacement; synthesis under the action of microwave radiation);
- rapid thermal decomposition of precursors in solution (spray drying; rapid expansion of supercritical fluid solutions; cryochemical);
- self-igniting (glycine-nitrate; Pechini method; cellulose (fabric, paper) technology; pyrolysis of polymersalt films).

The choice of using a particular method depends on the chemical nature of the obtained compounds, the size and morphological characteristics of the particles of the synthesis products, the conditions, and method of formation of the latter; the material and condition of the surface, the shape of the samples on which the coating is applied; capabilities of available technical equipment, etc. These methods are used both independently of each other and in combination.

Recently, titanium dioxide has attracted special attention due to new unique prospects for its use in the form of nanostructured materials and nanocomposites with controlled morphological, physicochemical and optical properties. TiO₂, which has high chemical and thermal stability, as well as impurity levels in the electronic structure of the material, created by a given type of doping, is unique for building on its basis new effective functional materials used in photocatalysis and photovoltaics, sensory, catalysis, for liquid chromatography and other areas.

The essence of the FC properties of TiO_2 is that in the volume of a semiconductor particle under the influence of electromagnetic radiation, electron - hole pairs are generated, which, upon reaching the surface of a TiO_2

particle, enter redox reactions with adsorbed molecules. For titanium dioxide, the process is as follows

$$(Ox_1)_{ads} + (Red_1)_{ads} \xrightarrow{TiO_2} Ox_2 + Red_2$$

In this case, part of the electrons and holes can undergo recombination in the bulk or on the surface of TiO_2 . For the effective occurrence of photocatalytic processes, it is necessary that redox reactions involving an electron-hole pair be more effective than recombination processes.

The advantages of the photocatalytic method of purification are well known: 1) the ability to oxidize almost any organic matter and also a number of inorganic, such as CO, H₂S, HCN, NH₃, NO_x, etc.; 2) the method works at room temperature and atmospheric pressure; 3) it is possible to oxidize even small concentrations of pollutants, cleaning of which by other methods is economically unprofitable; 4) for the implementation of the method of photocatalytic purification does not require additional reagents, because the oxidant is oxygen.

However, for the photocatalytic method of air purification, a number of disadvantages are known, such as:

- 1) a relatively low purification rate;
- 2) the need to use ultraviolet light sources in the case where the photocatalyst is titanium dioxide;
- 3) low adsorption capacity of most simple photocatalysts;
- 4) the possibility of formation of intermediate products during the oxidation of high concentrations of pollutants.

Therefore, the development of new photocatalytic systems that would overcome these shortcomings is the subject of an urgent series of modern studies.

Definition of unsolved aspects of the problem

When creating and heat treatment of materials based on titanium dioxide, it is necessary to take into account the possibility of surface and bulk defects of the TiO₂ crystal lattice (due to the existence of phase diversity in the range of O / Ti <2 ratios on the phase diagram of the Ti - O system [14]) and detection their significant effect on the photochemical properties and photoinduced hydrophilic ability of the synthesis products [15].

Available information on the state and possible directions of improvement of technologies for creating such materials, current requirements for their stability and reproducibility of properties, expansion of their use [1–15], manifestation of high activity of low-crystallized particles of structural components formed by solvent thermolysis [16], new information on reactivity and transformation of layered perovskite-like oxides, stabilization of photocatalytic and sensory-active crystalline modification of anatase due to NO₃⁻ ions [17], doping of Ln₂O₃ [18, 19] in obtaining TiO₂ from solutions [20] initiated the continuation of our study. Today, ways to control the technical parameters of the target products are through the choice of composition, synthesis conditions, and method of processing.

One of the most promising classes of complex oxide materials of rare earth elements and titanium are nanostructured layered perovskite-like compounds and solid solutions based on them. Depending on the composition and structure, they have a wide range of physicochemical properties. Presented in this paper perovskite-like layered titanates belong to the homologous series (Me, Ln)_{n+1} Ti_nO_{3n+1}, where Ln - La - Nd, Me - Li - Cs, n is the number of nanolayers of perovskite (Ruddlesden-Popper phases; with a thickness of approximately one layer 0.5 nm). Accordingly, MeLnTiO₄ in its structure contains one nanolayer of perovskite, Me₂Ln₂Ti₃O₁₀ - three. Perovskite-like nanolayered titanates were obtained and studied by the authors of [21- 24]. Figure 1 shows an extended unit cell for Me₂Ln₂Ti₃O₁₀ (Ln - La, Nd; Me - Li, K) [23] and Na₂Ln₂Ti₃O₁₀ [24].

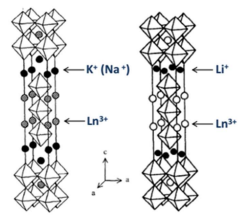


Figure 1 – Extended elementary cells of Ruddlesden-Popper phases, [23, 24]

So oxides of $K_2Nd_2Ti_3O_{10}$ (as an example) crystallize in a tetragonal structure. The spatial group for these compounds is defined as I4 / mmm. The thickness of layered oxides of this type is characterized by three titanium-oxygen octahedra $[Nd_2Ti_3O_{10}]$, alternating with each other and separated by alkali metal cations, in this case, potassium cations, between the layers. The lattice parameter $c \approx 30\text{\AA}$ indicates the displacement of adjacent perovskite layers by $\frac{1}{2}$. The neodymium cation is located in the center of the perovskite lattice and is characterized by a 12-coordinated oxygen environment. The alkali metal cation is located in the interlayer space and is usually 9-coordinated [25].

The layered structure, consisting of lamellar particles, is stored by the product when kept in humid air and confirmed by scanning electron microscopy [26]. The identified stability of the systems recommends them as promising photocatalysts in conditions close to their use and is important for other innovative areas of their applications.

Analysis of recent publications shows that titanium dioxide is mainly used in thin film form, which most effectively implements its properties necessary for photocatalysis, solar energy, sensors, self-cleaning coatings, and more.

And the practical implementation of modern technologies of the already proposed variant of the composite photocatalyst [27], the structure of the granules of which is represented by three layers: adsorbent, silicon

dioxide, and photocatalyst - titanium dioxide anatase modification, will simultaneously solve problems: 1) effective adsorption of both polar and nonpolar (for example, pollutants); 2) exclusion of the influence of the electrically conductive properties of the sorbent on the recombination of photogenerated electron-hole pairs; 3) to ensure complete absorption of incident light by the particles of the photocatalyst, not the adsorbent; 4) detection of photocatalytic activity under visible light.

Today, thanks to the technological methods of reactions of "soft chemistry", it is possible to create substances with various structural features, to obtain metastable compounds by a sequence of low-temperature topochemical syntheses. Such reactions with a change in the structure and morphology of the particles take place at low temperatures while maintaining the basic structural features in perovskite-like layered oxide compounds. Depending on the nature and stoichiometry of their cations, they can exhibit a variety of physical and chemical properties: superconductivity, colossal magnetoresistance, ferroelectricity, catalytic and photocatalytic activity, the ability to ion exchange in solutions and melts, and the ability to hydrate. Therefore, the study of the peculiarities of transformations of intermediate precursors - alkaline coordination nitrates of REE, their reactivity during the synthesis of layered perovskite-like oxide phases directly affects the possible areas of further application of the latter.

For perovskite-like compounds, such researchers include, in particular, ion exchange [25], intercalation and deintercalation [28], various substitution and condensation processes [29], fission processes [30], and mutual transformations of one structure into another [31] (for example, the transition from Ruddlesden-Popper phases to Dion-Jacobson phases; transition within one type of phase with increasing or decreasing number of layers).

The most common reactions of "soft" chemistry are ion exchange reactions, during which the weakly bound cations of the interlayer space are replaced, while the perovskite layers are quite stable mainly due to metaloxygen covalent bonds and play the role of a framework in the layered structure. This allows the substitution reactions of some interlayer cations to others, without affecting the basic structure of the layered oxide. Such reactions can be used to obtain a wide range of new perovskite-like structures.

For effective management of the properties of the received products, a deep understanding of physical and chemical processes, and the phenomena occurring during their formation are necessary. And their complex research with the application of modern physical and chemical methods allows improving our knowledge of characteristic features of fast-moving processes, stages of evolution of structure, and microstructure of technological objects.

The purpose and tasks of the study

To study the cooperative processes of interaction between structural components in the systems of nitrate precursors of REE representatives of the cerium subgroup and elements of IA subgroup of the periodic table (Li, Na, K) in conditions similar to the regulations for creating multifunctional oxide materials for various purposes using photocatalytically active TiO₂ to form reliable ideas and obtaining objective knowledge about the features of transformations and the total behavior of the constituent elements in the preparatory stages of processing of technological objects with thermal activation (25-1000 °C), necessary for improving and developing methodologies and regulations of modern manufacturing technologies.

Experiment Methodology

To evaluate the possibility of controlling the processes of multistage formation of complex oxide compositions with multifunctional properties and substantiation of phase formation mechanisms as a model using a set of physicochemical methods, water-salt nitrate systems MeNO₃ - Ln (NO₃)₃ - H₂O, (Me - Li, Na, K; Ln - La-Sm) at 25-100 °C. The choice of the composition of the objects of study, temperature sections are determined by a number of factors.

First, among the elements of the rare earth series, the highest complexing ability is found by representatives of the cerium subgroup; among them, the largest changes in the composition, structure, and properties of their compounds are the elements of its middle, Pr and Nd. The selected components of the systems specify the technical characteristics of the target product or are modifiers of its properties. And the presence of a large number for the use of potential electronic analogues (representatives of natural series of rare earth, alkaline elements) causes significant variability and breadth of the range of modification of their characteristics. Temperature cross sections are due to the areas of the existence of crystal hydrate forms of the source components.

Secondly, according to [32], three layered potassium titanates $K_2Ln_2Ti_3O_{10}$ (Ln - La, Nd) of the above components, obtained by ceramic technology, in suspension form in aqueous-alcoholic solutions under the action of UV radiation show the highest among known species phases photocatalytic activity for the decomposition of H_2O and the release of hydrogen. This fact is associated with the morphology of their particles and the ability to reversibly intercalate water molecules in the interlayer space, which can lead to both an increase in the effective specific surface area of the photocatalyst and contribute to the spatial distribution of redox centers.

To determine the nature of chemical interaction and phase equilibria in water-salt systems of the studied nitrates (precursors of multicomponent oxide polyfunctional materials) in full concentration ratios in the temperature range of solutions using the method of additives described in [33, 34] and based on the study of solutions one of the properties of the most "sensitive" to the detection of phase transformations in systems, which is both a parameter of their state, and also simple experimental methods currently available. The method allows to find the limits of self-development to which the isolated system of the set structure goes in concrete conditions in an equilibrium state.

The equilibrium of the phases was reached within 2-3 days. Hydrated and anhydrous nitrates of these "c.f.a". elements were used as starting salts.

Chemical analysis of liquid and solid phases, "residues" was performed on the content of Ln^{3+} and nitrogen. The Ln^{3+} content was determined trilonometrically in the presence of xylenol orange as an indicator (acetate buffer solution, pH = 5–6) [35]; nitrogen - by distillation; Me⁺ ions - calculated by the difference, based on the total nitrate content and partly on the dry residue.

The obtained data for individual ions were converted into salt content and plotted on the solubility diagrams according to the principle of conformity. Graphical representation of the composition of solid phases formed in the system was performed according to Screenmakers [33, 34], confirmation of their individuality and characterization - chemical, crystal-optical, X-ray phase, X-ray structural, IR spectroscopic, thermographic, and other methods.

Crystal-optical determinations of the compounds were performed by the immersion method using a MIN-8 microscope. Phase analysis was performed on a DRON-3M diffractometer (Cu Kα radiation, Ni filter) by the "powder" method. Diffractograms were deciphered from the PDF file JCPDS. Determination of symmetry, parameters of unit cells, and measurement of the intensity of diffraction reflections from single crystals was performed on an automatic X-ray single crystal diffractometer CAD - 4F "Enraf - Nonius" (Mo $K\alpha$ - radiation, graphite monochromator; ω / 2θ method). All calculations for the determination and refinement of atomic structures were performed using complexes of crystallographic programs SHELX, XTL - SM, AREN. The IR absorption spectra of the synthesized compounds in the region of 400–4000 cm⁻¹ were recorded on a UR - 20 spectrophotometer using a standard vaseline oil suspension technique. Thermograviometric analysis was performed on a Q – 1500 D derivatograph at temperatures from 293 K to 1273 K in the air with a heating rate of 10 degrees. / min and the developed device for DTA.

Basic material and results

Generalized and important for practical use information on alkaline coordination nitrates of rare earth elements of the cerium subgroup - precursors of promising modern multifunctional materials - on the conditions of their formation and existence, nature of the chemical bond, composition, structure, type of ligand coordination, the existence of isotype series on stoichiometry, composition the structures of the revealed properties are systematized by authors and in the most obvious form have resulted in tab. 1-4. The choice of this form of data presentation is the most informative and useful in the development of innovative projects that allows predicting the causal fundamental patterns of behavior of structural components in similar production processes, to choose the right modes, stages, methods of forming, and obtaining target products with reproducible structurally sensitive characteristics.

Table 1 – Isothermal concentration limits of crystallization of alkaline coordination nitrates of neodymium from solutions of water-salt systems $MeNO_3 - Nd(NO_3)_3 - H_2O$ (Me – Li, Na, K)

| The composition of the compounds | t, °C | Compositions of responding to poi | The nature of solubility | |
|---|-------|-----------------------------------|--------------------------|-------------|
| | | MeNO ₃ | $Nd(NO_3)_3$ | |
| $\text{Li}_3[\text{Nd}_2(\text{NO}_3)_9] \cdot 3\text{H}_2\text{O}$ | 65 | 19,32 | 59,61 | incongruent |
| | | 13,95 | 65,63 | |
| | 100 | 24,03 | 54,68 | congruent |
| | | 9,68 | 72,51 | _ |
| $Na_2[Nd(NO_3)_5]\cdot H_2O$ | 50 | 16,55 | 51,62 | incongruent |
| | | 8,60 | 62,58 | |
| | 65 | 20,44 | 50,40 | incongruent |
| | | 3,03 | 70,17 | |
| | 100 | 25,27 | 47,28 | incongruent |
| | | 4,15 | 76,96 | |
| K ₂ [Nd(NO ₃) ₅ (H ₂ O) ₂] | 50 | 27,26 | 51,62 | incongruent |
| | | 21,34 | 54,91 | |
| $K_3[Nd_2(NO_3)_9]\cdot H_2O$ | 50 | 21,34 | 54,91 | incongruent |
| | | 11,49 | 63,31 | |
| | 65 | 32,57 | 47,88 | congruent |
| | | 8,67 | 70,44 | |
| | 100 | 40,15 | 45,02 | congruent |
| | | 3,39 | 76,44 | |

The revealed regularities in the nature of the behavior of structural components in rubidium, cesium nitrate systems La - Sm, in objects based on REE of the yttrium subgroup (Y, Gd - Lu) indicate the possibility of only limited or special application of the latter predecessors in the study area. There are a number of objective and economic reasons for this. These are the features of the electronic structure of their atoms, lower manifestation of chemical activity and complex-forming ability of these Ln3+ in comparison with the elements of the cerium subgroup, weaker effect of the considered influencing factors on the studied processes. To clarify the general patterns and build a holistic objective picture of the behavior of such technological precursors, the authors studied the systems of natural series Y, La - Lu, Li - Cs. Analysis of the results of the study was published in previous works of the authors [10, 11, 36].

In ternary REE-containing nitrate precursor systems, which are integral components of more complex multicomponent systems, exchange transformations begin from the moment the components are dissolved in water. It was found that the Ln³+ cerium subgroup in the studied conditions is active complexing agents, that form anionic coordination compounds of Me+ of all alkali metals, and their stability and complex of inherent properties are potent technological factors that significantly affect the nature of transformations in systems as intermediates. and the results of processes in general.

The obtained information allows modeling the behavior of structural components at the preparatory stages of the formation of modern multifunctional photocatalytically active materials according to innovative technological regulations with the use of nitrate REE-containing precursors.

The available identified trends in phase formation in the model systems studied are thermodynamically the most probable limits of transformations in technological objects in the conditions of formation and production of target products. And possible real deviations in such systems are caused by inhomogeneity of the reaction medium in composition, the content of reacting components, conditions of their location, finiteness of transformations, diffusion features, heat capacity, viscosity, nature of transformations at the boundaries of formed hetero phases, using their applied principles and methods. activation and other specific factors. And the identified processes of complexation in aqueous solutions of nitrates contribute to the homogenization of systems of structural components at the molecular level in complex or combined processing.

The analysis of the obtained data indicates the flow of competing processes of replacement of water molecules by nitrate ions in the systems in the immediate vicinity of the Ln3+ complexing agent. The degree of completeness of substitution depends on the nature of Ln³⁺, the presence of Me⁺, the properties of electrondonor oxygen atoms and the spatial structure of ligands, the concentration of anions, the amount of solvent. A significant influence of the temperature factor on these processes is revealed. There are differences in the complexing ability of the elements of cerium and yttrium subgroups, Y, as well as among REE in the middle of the first subgroup. The obtained results indicate the gradual processes of complexation. The presence of certain values of the temperature at the beginning of the release into the solid phase of complex compounds - the existence of an energy barrier and the need for some activation energy to carry out such transformations. In the formation of nitrate complexes, the requirements

of symmetry are largely met, and the planar small-sized ligand NO₃⁻ is "convenient" for the formation of a highly symmetric environment of the Ln₃⁺ ion. Lanthanides have a tendency to form three types of NO₃⁻ ligand coordination. This leads to the formation of both isolated complexes and their polymerization into dual-core, chains, and frameworks.

All established coordination compounds were synthesized in single crystal form and characterized by a set of physicochemical methods. Table 2 shows the X-ray diffraction characteristics of the newly formed phases for the possibility of their identification and detection during processing.

Table 2 – Radiographic data of neodymium lithium, sodium, potassium coordination nitrates

| Li | $\text{Li}_3[\text{Nd}_2(\text{NO}_3)_9] \cdot 3\text{H}_2\text{O}$ $\text{Na}_2[\text{NO}_3] \cdot 3\text{H}_2$ | | Na ₂ [Nd(1 | $[Nd(NO_3)_5] \cdot H_2O$ $K_2[Nd(NO_3)_5 (H_2O)_2]$ | | | | K ₃ [Nd ₂ (NO ₃) ₉]·H ₂ O | | | | | |
|-------|--|-------|-------------------------|--|----------------------|-------|-------------------------|--|-------------------------|-------|-------------------------|-------|----------------------|
| d, Å | I/I ₀ ,% | d, Å | I/I ₀ , % | d, Å | I/I ₀ , % | d, Å | I/I ₀ , % | d, Å | I/I ₀ , % | d, Å | I/I ₀ , % | d, Å | I/I ₀ , % |
| 8,36 | 30 | 2,013 | 19 | 7,84 | 63 | 5,42 | 66 | 2,056 | 24 | 9,48 | 85 | 2,281 | 21 |
| 7,64 | 77 | 1,979 | 19 | 7,54 | 47 | 5,27 | 90 | 1,993 | 27 | 7,74 | 92 | 2,249 | 69 |
| 6,68 | 13 | 1,944 | 28 | 7,07 | 73 | 4,94 | 45 | 1,947 | 26 | 7,65 | 32 | 2,189 | 54 |
| 6,00 | 38 | 1,931 | 26 | 5,18 | 27 | 4,11 | 72 | 1,777 | 11 | 5,36 | 39 | 2,108 | 68 |
| 5,75 | 79 | 1,855 | 15 | 4,23 | 100 | 3,88 | 15 | - | - | 5,27 | 40 | 2,082 | 37 |
| 5,42 | 98 | 1,778 | 15 | 3,80 | 57 | 3,80 | 42 | _ | - | 4,94 | 30 | 2,058 | 15 |
| 5,26 | 74 | 1,726 | 28 | 3,15 | 23 | 3,66 | 12 | - | - | 4,76 | 46 | 2,012 | 49 |
| 4,76 | 51 | 1,708 | 34 | 3,09 | 67 | 3,53 | 42 | _ | - | 4,49 | 87 | 1,909 | 21 |
| 4,64 | 100 | _ | _ | 3,02 | 86 | 3,35 | 38 | - | - | 4,26 | 27 | 1,837 | 27 |
| 4,35 | 43 | - | _ | 2,629 | 20 | 3,18 | 13 | - | - | 4,06 | 100 | 1,757 | 19 |
| 4,19 | 47 | - | _ | 2,391 | 17 | 3,05 | 73 | _ | - | 3,89 | 40 | 1,729 | 16 |
| 3,94 | 51 | _ | _ | 2,346 | 17 | 2,873 | 17 | _ | - | 3,78 | 22 | 1,714 | 18 |
| 3,90 | 40 | _ | _ | 2,307 | 8 | 2,843 | 33 | _ | - | 3,73 | 39 | _ | _ |
| 3,56 | 26 | _ | _ | 2,234 | 13 | 2,783 | 14 | - | - | 3,36 | 19 | _ | - |
| 3,32 | 19 | _ | _ | 2,178 | 27 | 2,750 | 17 | - | - | 3,27 | 26 | _ | - |
| 3,22 | 47 | _ | _ | 1,979 | 13 | 2,724 | 40 | _ | - | 3,18 | 55 | _ | _ |
| 2,978 | 34 | _ | _ | - | _ | 2,664 | 14 | - | - | 3,07 | 16 | _ | - |
| 2,772 | 19 | - | _ | - | _ | 2,639 | 19 | _ | - | 3,04 | 17 | _ | _ |
| 2,617 | 28 | - | _ | - | _ | 2,594 | 100 | - | - | 2,844 | 20 | _ | - |
| 2,545 | 43 | _ | _ | - | _ | 2,463 | 37 | - | - | 2,755 | 12 | _ | - |
| 2,385 | 21 | _ | _ | _ | _ | 2,392 | 19 | _ | - | 2,730 | 18 | _ | - |
| 2,328 | 17 | _ | _ | _ | _ | 2,374 | 44 | _ | - | 2,647 | 55 | _ | _ |
| 2,305 | 23 | _ | _ | _ | _ | 2,314 | 15 | _ | - | 2,592 | 16 | _ | - |
| 2,226 | 19 | - | _ | - | _ | 2,235 | 25 | - | - | 2,508 | 87 | _ | - |
| 2,135 | 49 | _ | _ | _ | _ | 2,188 | 10 | _ | - | 2,468 | 78 | _ | _ |
| 2,111 | 43 | _ | - | - | ı | 2,099 | 21 | - | - | 2,349 | 32 | _ | _ |

Note: d, Å - interplanar distances; I/I₀, % - the relative intensities of the reflexes.

The authors conducted a crystal chemical analysis of alkaline rare earth nitrate compounds, which is based on the results of our own research [37, 11] and literature data [38]. Particular attention is paid to the structure of coordination polyhedra Ln, which largely determines the basic properties of the corresponding compounds. Analyzing the shape, symmetry, and other properties of polyhedra (see Table 3), we get the opportunity to approach the understanding of the individual features of REE.

Data on average Ln - O distances in rare earth nitrates of alkaline cations discussed in this paper are in good agreement with the expected tendency to decrease Ln - O distances according to lanthanide compression and to increase these distances with increasing coordination number for fixed REE ion. The distances Ln - O (H₂O), as a rule, are among the shortest contacts in polyhedra. This fact can be explained on the basis of the presence of competing interactions with bonds of the type Ln - O (NO₃-). Coordination polyhedra are composed, as a

rule, of oxygen atoms of bidentate coordinated nitrate groups, less often in combination with oxygen atoms of water molecules. In all considered compounds coordination numbers of representatives of lanthanides of cerium subgroup 12.

When summarizing the structural data of alkaline coordination nitrates, REE draws attention to the extremely limited range of coordination polyhedra for such high CN REEs. In Fig. 2 shows a general view of he identified types of icosahedrons found in the structures of the studied compounds. In cases where water is not part of the coordination sphere of the Ln atom, the polyhedra are constructed in exactly the same way (Fig. 2a). Their equivalence is that the shortened O – O ribs (common to NO₃- - ligands) occupy the same positions. Exceptions are one of the 2 independent polyhedra in two isostructural compounds $\text{Li}_3[\text{Ln}_2(\text{NO}_3)_9] \cdot 3\text{H}_2\text{O}$ (Ln - La, Nd) (Fig. 2b). If the coordination saturation occurs with the participation of water molecules, the shape of the 12-vertices changes

slightly. Topologically, it is still the same icosahedron. However, the distribution of shortened ribs in this case is different (Fig. 2c).

The analysis of the Ln-polyhedra considered by us gives grounds to note the tendency of REE in this class of compounds to the organization of a symmetric coordination environment. In some cases, despite the fact that Ln³⁺ ions are located in common positions, their coordination polyhedra have at least one noncrystallographic axis of symmetry 2 (see Table 3).

The following data from the thermographic study of lithium, sodium, potassium coordination nitrates of rare earth elements of the cerium subgroup (Fig. 3, Table 4) clarify the nature and patterns of thermal transformations of these compounds in the temperature range 25-1000 °C, establish their heat resistance, intervals, staged processes, phase formation depending on the composition, content, nature of the components, the method of packing coordination polyhedra in spatial

construction, conditions and method of processing, etc. This makes it possible to predict the behavior of the applied potential predecessors in real technological systems in similar conditions. For comparison and analysis there is information about the low stability and heat resistance of potassium coordination nitrates Y, Gd - Lu composition $K[Ln(NO_3)_4(H_2O)_2]$, which leads to limitations in the use of this type of precursors in technological transformations to modify the properties target products.

It was found that the coordination numbers of the Ln³⁺ cerium subgroup - 12, found in low-temperature associated forms, remain unchanged until the formation of stable high-temperature multicomponent oxide phases MeLnO₂, Me₂Ln₂Ti₃O₁₀. This indicates the feasibility of using alkaline coordination nitrate REE-containing precursors in such technological innovative solutions.

| Table 3 – Coordination polyhedra of Ln atoms in alkaline rare earth nitrates |
|--|
|--|

| Compound | D_1 | D_2 | D_3 | C. N | Polyhedron |
|--|-------|-------|-------|------|------------------------------------|
| | | | | | type, |
| | | | | | its symmetry |
| Li ₃ [La ₂ (NO ₃) ₉]·3H ₂ O | | 2,66 | 2,66 | 12 | Ic 4L ₃ 3L ₂ |
| | | 2,65 | 2,65 | 12 | Ic L ₃ 3L ₂ |
| $\text{Li}_3[\text{Nd}_2(\text{NO}_3)_9] \cdot 3\text{H}_2\text{O}$ | | 2,61 | 2,61 | 12 | Ic 4L ₃ 3L ₂ |
| | | 2,60 | 2,60 | 12 | Ic L_33L_2 |
| $Na_2[Nd(NO_3)_5]\cdot H_2O$ | | 2,62 | 2,62 | 12 | Ic L ₃ 3L ₂ |
| | | 2,61 | 2,61 | 12 | Ic L_33L_2 |
| | | 2,61 | 2,61 | 12 | Ic L_33L_2 |
| K ₂ [La(NO ₃) ₅ (H ₂ O) ₂] | 2,70 | 2,68 | 2,68 | 12 | Ic L ₂ |
| $K_2[Nd(NO_3)_5 (H_2O)_2]$ | 2,62 | 2,61 | 2,61 | 12 | Ic L ₂ |
| $K_3[Pr_2(NO_3)_9]$ | | 2,63 | 2,63 | 12 | Ic L ₃ 3L ₂ |
| $K_3[Nd_2(NO_3)_9]$ | | 2,61 | 2,61 | 12 | Ic L ₃ 3L ₂ |
| (NH ₄) ₂ [La(NO ₃) ₅ (H ₂ O) ₂]·H ₂ O | 2,54 | 2,70 | 2,67 | 12 | Ic L ₃ 5L ₂ |
| $(NH_4)_2[La(NO_3)_5 (H_2O)_2] \cdot 2H_2O$ | 2,59 | 2,70 | 2,68 | 12 | Ic L_35L_2 |
| (NH ₄) ₂ [Pr(NO ₃) ₅ (H ₂ O) ₂]·2H ₂ O | 2,54 | 2,66 | 2,64 | 12 | Ic L ₃ 5L ₂ |

D₁, D₂, D₃ – average distance (Å) 〈Ln–O _{H2O}〉, 〈Ln–O _{NO3}〉, 〈Ln–O〉; Ic – icosahedron

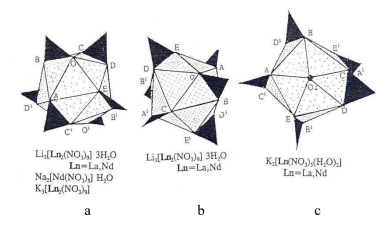
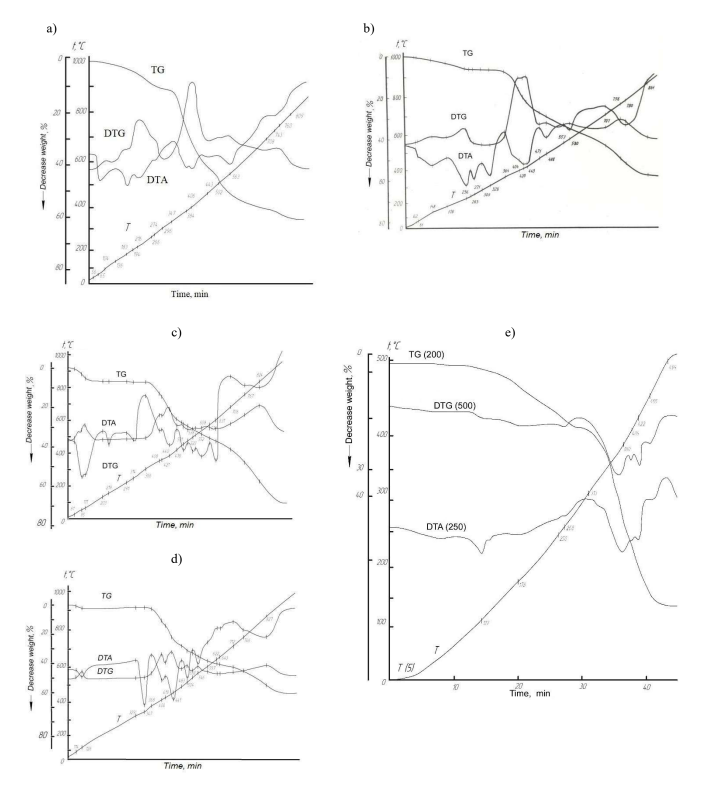


Figure 2 – Schematic representation and general view of Ln-icosahedrons found in the structures of the REEs of the cerium subgroup and lithium, sodium, and potassium



 $Figure 3 - Derivative diagrams: \\ a - Li_3[Nd_2(NO_3)_9] \cdot 3H_2O; \ b - Na_2[Nd(NO_3)_5] \cdot H_2O; \\ c - K_2[Nd(NO_3)_5 (H_2O)_2]; \ d - K_3[Nd_2(NO_3)_9] \cdot H_2O; \\ e - K[Yb(NO_3)_4(H_2O)_2] (Ln - Y, Gd-Lu) \\ T - temperature curve; DTA - differential temperature curve; \\ \label{eq:definition}$

TG – thermogravimetric curve; DTG – differential thermogravimetric curve

Table 4 – The temperature values of the detected effects during the heat treatment of the representatives of the established groups of alkali coordination nitrates REE

| Compounds; spatial group of crystals | Representatives | Temperature interval of formation, °C | The nature of solubility | Dehydration | Melting in crystallization water | Polymorphic transitions | Melting anhydrous form | The composition of the products of conversion at $980^{\circ}\mathrm{C}$ |
|---|-----------------|---------------------------------------|--------------------------|------------------|----------------------------------|-------------------------|------------------------|--|
| Li ₃ [Ln ₂ (NO ₃) ₉]·3H ₂ O cubic.; P2 ₁ 3 | La –Sm | 65-100 | congr. | 65 183 216 | 183 | - | 274 | LiLnO ₂ |
| Na ₂ [Ln(NO ₃) ₅]·H ₂ O monocl.; P2 ₁ /a | La –Sm | 50-100 | congr. | 81 148 236 | _ | 271 | 328 | NaLnO ₂ |
| K ₂ [Ln(NO ₃) ₅ (H ₂ O) ₂] rhomb.; Fdd2 | La – Nd | 50-100 | incongr. | 95,111 | 95 | 219 | 314 | KLnO ₂ , Ln ₂ O ₃ |
| K ₃ [Ln ₂ (NO ₃) ₉]·H ₂ O cubic.; P4 ₃ 32 | La –Sm | 50 | congr. | 126 | _ | - | 347 | Ln ₂ O ₃ |
| K[Ln(NO ₃) ₄ (H ₂ O) ₂] prim. rhomb.; P2 ₁ cn | Y, Gd - Lu | 50-100 | congr. | 138, 172 | 138 | = | = | Ln ₂ O ₃ |

Scientific novelty

A comprehensive systematic study of the interaction of structural components in systems of rare earth nitrates and IA groups of elements of the periodic table precursors of modern multicomponent oxide functional materials based on them - established the formation of a wide class of alkaline coordination nitrates of lanthanides. The identified objective patterns have a fundamental and applied value, and deepen the understanding of:

- the chemical and physical properties of Ln, their complexing ability,
- the possibility of formation and existence in similar systems of associated new phases, their atomic-crystalline structure, stability, and stability,
- the influence of the nature of lanthanides and alkali metals on the structure of complex anions and compounds in general,
- the individuality of Ln complexes,
- the existence of isotypic in composition and structure of groups of compounds in the natural series of lanthanides and alkali metals,
- the role of NO₃⁻ groups in the stereochemistry of this class of nitrates,
- the role of water in the formation of the immediate environment of Ln³⁺ ions-complexing agents.

Practical significance

The obtained system of knowledge about transformation processes in REE-containing nitrate precursor systems and crystal chemical properties of Ln coordination nitrate samples acquires special value in the formation of nanostructured layered perovskite-like compounds of lanthanides and transition elements (including titanium, described in [7, 12, 32, 39, 40], others), solid solutions based on them; in establishing technological and functional dependencies between the method of preparation, variability of the method of activation of technological systems, methodology of manufacturing the target product and its phase composition, lattice parameters, specific surface area, the morphology of constituent particles, the activity of selfcleaning compositions with photocatalytic and hydrophilic properties, and special structural elements; in the practical implementation of innovative projects of water decomposition for the purposes of hydrogen production (as an alternative fuel), decomposition of toxic organic substances in solutions and air, incomplete oxidation of carbohydrates; in obtaining other perovskitelike phases by ion exchange reactions and in other areas.

Conclusions

- 1. The results of the study show that the processes of obtaining oxide REE-containing structural and functional materials for different purposes using nitrates of elements of different electronic structures by chemical mixing of the source components in the joint separation of products from the liquid phase by sequential or coprecipitation followed by heat treatment. Data on their composition, content, and behavior in each case require prior systemic empirical knowledge in full concentration ratios in a given temperature range.
- 2. Differences in the behavior of structural components in the systems of lanthanides of cerium and yttrium subgroups, in their nature of the interaction, stages, features, and patterns of flow.

- 3. The new knowledge is the basis for:
- finding ways to increase the activity of Ln-forms;
- elucidation of the nature of successive thermal transformations in nitrate REE-containing multicomponent systems of different aggregate states during their heat treatment; conditions of formation and existence, properties of intermediate phases; influencing factors; possible ways to control the receipt of the target product;
- in case of creation of modern perfect low-cost technologies of formation of functional materials of various functions with reproducible properties.

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