Research Article

Production of Nanopowders of Metal Oxides Using Pulsed Electron Beam in Low Pressure Gas

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The installation for production of metal oxide nanopowders was created. The method involves evaporation of the target by a pulsed-electron beam, condensation of the vapors of the material in a low-pressure gas, and deposition of nanopowders on a cold large-area crystallizer. In a new installation, a higher-power electron gun with a hollow cathode, which ensures the formation of the current pulse of the electron beam with amplitude up to 1 A and a duration of 100 μs, and a crystallizer of a larger diameter (0.3 m) and length (0.5 m), which makes it possible to decrease the agglomeration, were used. The results of the evaporation of targets made of YSZ, CeGdOx, Zn-ZnO, Al2O3, and ZnO were presented. Room-temperature ferromagnetism has been observed in YSZ, Zn-ZnO, and Cu(Al)-doped Al2O3. The proposed method makes it possible to obtain nanopowders of oxides with a characteristic particle size of 3–5 nm and agglomerates consisting of them 20–600 nm in size, specific surfaces of up 338 m2/g, productivity of up to 12 g/h, and a specific power consumption ≥112 (W h)/g.

1. Introduction

Production of weak aggregated nanopowders (NPs) with the characteristic size in the field of less than 10 nm and narrow distribution of particles in the sizes remains is an actual problem, especially, for complex chemical compounds at high cleanliness of a received product. Thus it is desirable, that the method provided the possibility to use as raw materials enough cheap substances and probably wider range of it (conducting and nonconducting, alloys and metals, mechanical mixes, etc.).

For many problems ways of producing NPs by a method of pulse heating and target evaporation passing a pulsed current-electric explosion of wire (EEW) [1] and pulsed CO2 laser irradiation [2] is successfully used. EEW allows us to produce NP metals and alloys, and also their chemical compounds at rather small consumed energy (<50 W h/g) and high efficiency (50–400 g/h). However, produced NPs have rather wide size distribution of particles and it is not possible to prepare all desirable metals and alloys in the form of a metal wire of certain diameter and structure.

For production of NP by target evaporation irradiation of various types of lasers is used, but the greatest application have found CO2 [2] and fiber lasers [3]. Using of the continual CO2 laser, it has been shown [4, 5] that productivity of process depends on capacity of radiation and its intensity, productivity reaches 130 g/h at consumed energy 25 W h/g, and the size of particles \( d_{BET} = 60 \) nm for ZrO2 powders.

The pulse mode should provide reduction of the size of particles, at invariable consumption of energy, at the expense of the best removal of particles from a hot zone and faster decrease in concentration of steams, and also possibility of increase intensity of radiation. For realization of a pulse mode of evaporation was used pulse CO2 [5, 6] which allows to reduce essentially width of distribution of received particles till the size, and also to use as raw materials for targets cheap large powders. At equal power consumption, in comparison with the continual CO2 laser, use of pulse irradiation has allowed to lower the characteristic size of particles from 4 to 6 times. It was experimentally shown that the major factor defining productivity of installation with set characteristics, was specific energy of evaporation of
a material. It is important to notice that the method provides preparation of pure NPs and is ecologically pure.

However, this method has small efficiency (1-2%) in the use of energy because of its losses at radiation generation, and also at interaction of radiation with a plasma jet and substance. Productivity of a method was from 15 to 80 g/h and, basically, was defined by the energy which is required for evaporation powders of chosen structure. The basic problems connected with insufficient efficiency and small operating time CO2-laser was eliminated at transition to fiber lasers [7].

In our point of view to increase the efficiency of energy transformation in radiation and use the irradiation for the evaporation, heating a target by an electron beam looks more attractive. Unlike laser radiation, using the electron beam, being formed on a target surface plasma absorbs arriving energy of the beam according to density of steam of target material. Thus, target heating under a vapour-plasma jet proceeds, though and with slightly smaller capacity. At use of accelerators with high electron energy (>1 MeV) to evaporation of targets in high-pressure gas that simplifies questions of cooling of a steam phase and collection of the formed powder [8] was possible. But such accelerators with necessary radiating shield are expensive enough.

Electron guns with energy to 50 keV are rather cheaper; however in this case the beam injection to atmosphere is impossible. The available literary data on use of such guns with continuous beam [9–11] for production NPs simple oxides at low pressure of gas has shown that in all researches powders oxides with the size of particles less than 5 nm were produced. However in work [9] the particles was poorly agglomerated and consumed energy was 200 W h/g for Al2O3 whereas in [10, 11] powders only in the form of agglomerates of particles were produced, and consumed energy have increased to 1000 W h/g and more 4000 W h/g [11].

These contradictions, desire to lower an energy expense and also to investigate production possibilities of NPs with complex structures stimulated us to create the installation with a pulsed electron gun and to the beginning of experiments in this direction.

It is necessary to consider that at heating by an electron beam excluding the creation of a liquid bath completely was impossible, as absorption of energy of an electron beam has nonlinear distribution [12]. Expansion in gas of low pressure allows receiving high speeds of decrease in concentration of steams that promotes production of small sizes particle, but speed of cooling of steams and particles that allow their aggregation simultaneously falls. A doubtless advantage of pulsed electron beam (PEB) is that it is possible rather simply to change, except electrons energy, a pulse width and repetition rate that expands experimental possibilities. Besides, to operate position of PEB it is simple enough at the expense of its scanning that facilitates a problem of uniform evaporation of a target, in difference from lasers.

The first installation “Nanobim-1” for production nanopowder by a method of PEB evaporation of target [13], using a hollow cathode gun [14], with condensation of steams of a material in low-pressure gas and condensation NPs on a cold crystallizer, has been placed in KAERI (Republic Korea) and has shown a realizability and certain advantages and disadvantages of the method [13]. The received experience has allowed us to create the modernized installation “Nanobeam-2” using which various NPs have been received and their properties were investigated.

2. The Description of Installation and Experiments Conditions

Installation diagram of the “Nanobeam-2” is given in Figure 1, and its technical characteristics are in Table 1. For normal work of the chosen electron gun pressure about 10⁻² Pa is required, and for fast cooling of particles it is desirable to have a high pressure in the evaporation chamber. Calculations on [15] for the chosen geometry of the chamber of drift (length 20 cm), have shown that at reasonable losses of energy of a beam (not more than 10%) in the chamber of drift the pressure size in the evaporation chamber can be 100 Pa.

These inconsistent requirements are provided with pumping out from three zones divided two gaz-dynamic windows [17] with apertures in diameter of 3.2 mm, 100 mm placed on distance from each other in the drift chamber where focusing and rejecting coils are located also.

Pumping out of air from chambers of an electron gun and drift was carried out by diffusion pumps by productivity 160 and 100 l/sec, accordingly, and from the evaporation chamber-mechanical pump productivity 60 l/sec.

All chambers were made of stainless steel, by sheet lead in the thickness of 5 mm for X-ray radiation shield are outside covered, viewing windows were protected by lead glass in the thickness of 20 mm.

Operating experience of installation “Nanobeam” has shown that the electron gun [14] forms insufficient amplitude of a current and demanded energy was necessary to type at the expense of increase of pulse width that raised losses of energy at the expense of heat conductivity. Therefore the new design of an electron gun with the plasma emitter which generates a pulse electron beam with frequency of repetition of 1–500 Hz and duration of impulses of a current 100 μsec, at accelerating voltage up to 40 kV with a current about 1 A [18] has been created.

Beam scanning on a target is carried out continuously by a principle of development of a TV camera, thus the scanning area could change from 0.8 to 4 cm².

Target (a tablet in diameter of 40–60 mm and height to 20 mm, Figure 2) it is established in a support having free rotation (with fixing) in two axes.

At an adjustment and focusing electron beam the target is established horizontally. For evaporation the target turns at an angle 45° to the beam in the crystallizer direction for increase in collecting the powder. The support design allowed to rotate a target with speed of 8,7 turns/min, for its more uniform evaporation, and also to replace it from a cut of the rejecting coil in limits from 80 to 170 mm.

Used in [13] the powder collection system on the basis of a rotating cooled hollow disk it has appeared difficult in operation. Besides, at increase in distance from a target
the share of a collected powder essentially decreases to a disk for the geometrical reasons. The modernized powder collection system is executed on the basis of the hollow copper cylinder, with internal diameter 300 mm which is established perpendicularly to a beam axis, on distance of 7 mm from target edge. The cylinder is cooled from within by liquid nitrogen with the expense about 5 kg/hour. From the cylinder the powder collected manually using a scraper. For simplification the technology of collection and control of doping NPs the cylinder surface can be covered by sheet from other metal (usually stainless steel).

The following methods were used for examination of the materials.

(1) The specific surface of the powders $S_{BET}$ was measured by the BET method on a TriStar 3000 V6.03 analyzer.

(2) The chemical composition of the powders was determined and the X-ray diffraction analysis (XRD) was performed using standard methods on a Discover D8 X-ray diffractometer.

(3) The microscopic analysis was made in electronic microscopes SEM LEO-982 and TEM JEOL 2100.

Phase transformations, the concentration, and the absorption of gases in the powders were determined by the methods of scanning differential calorimetry (DSC) and thermogravimetric analysis (TG) on an STA 409 PC Luxx.
thermal analyzer and a QMS 403 Aéolos mass spectrometer (NETZSCH).

The chemical analysis were carried out by the method of inductive and connected plasma (ICP) on the spectrometer iCAP 6300 Duo.

Magnetic characteristics NPs defined on the scales of Faraday.

The accelerating voltage of 37–40 kV was used in the experiments. Upon transportation and focusing the beam current on the target was not less than 0.46 A at the beam diameter of about 1.0 mm, providing the power density (the intensity) of about $10^6$ W/cm².

The pressure of the evaporation chamber was adjusted between 3.8 and 50 Pa. The beam energy loss caused by this adjustment was several percent and was disregarded. Air, oxygen, and argon were used in the experiments and the gas inleakage rate was controlled by rotameters.

The total current of the electron beam and the beam current hitting the target were measured by current transformers installed in the accelerating voltage source and on the target support acting as a current lead. The shunt in the connection between the upper gas dynamic window and the casing allowed measuring the proportion of the current lost as the beam was transported through the window.

The operating principle of the device is as follows. The electron beam is focused at the hole of the upper gas dynamic window and, as it passes through the second gas dynamic window, it focused additionally on the deflecting coil. This coil also sweeps the beam on the target. The material of the target evaporates under the action of the beam; the vapor-plasma mixture is cooled by the low-pressure gas in the evaporation chamber; and the vapor condenses with the formation of nanopowder particles. The powder flies to the cooled crystallizer and is deposited thereon.

3. Results of Experiments

Installation characteristics were defined in experiments on production of powders various oxides. Targets were carried out from industrial powders of demanded oxides in the micron sizes, pure and with various alloying additives, by the method of manual pressing with the subsequent roasting, for receiving ceramics, because not annealed targets collapsed from influence of IEP and did not allow to receive NPs with the high specific surface $S_{BET}$.

The synthesized YSZ (zirconia stabilized with 10 molar % of yttria) powders had a large-specific surface and were agglomerated to complexes of 20 to 600 nm in size, which consisted of particles with a sufficiently narrow size distribution of about several nm (Figure 3). The specific surface of the synthesized YSZ powders could be as large as $S_{BET} = 250$ m²/g and after sedimentation to $S_{BET} = 270$ m²/g.

XRD of the synthesized powder showed that the sample had two phases.

(i) The first phase was a solid solution of $Y_2O_3$ in the cubic modification of $ZrO_2$. The lattice spacing $a = 5.148$ Å and the coherent scattering regions (CSR) were 130 nm in size. The $Y_2O_3$ concentration was $\approx 11$ mole %. The varying concentration of yttria suggested that the phase was not fragments of the target, but represented a synthesis product. The concentration of this phase was $\approx 7$ weight %.

(ii) The second phase ($\approx 93$ weight %) was an X-ray amorphous phase with the short-range order characteristic of the monoclinic form of zirconia having the following parameters: $a = 5.17$ Å, $b = 5.10$ Å, $c = 5.44$ Å, and $\beta = 86.9^\circ$. The CSR was $1 \pm 2$ Å in size on the average.

The obtained result is nontrivial. Earlier investigations into synthesis of zirconia nanopowders (our data and the literature [19, 20]) showed that the tetragonal and cubic phases, which are most metastable at room temperature, are stabilized with decreasing size of the particles. In this experiment the finest phase is the monoclinic one, that is, the most stable phase. This would not be so if pure zirconia is synthesized. We synthesized yttrium-stabilized $ZrO_2$. 
The stable phase for this compound is the cubic modification. The most metastable is the monoclinic phase and, naturally, it is this phase that forms in finest particles. For understanding of real distribution of a yttrium on particles depending on their size of research it is necessary to continue.

Results TG-DSC of the analysis have shown (Figure 4) that at temperature nearby 187°C on curve DSC is observed strong exothermic peak caused by crystallization of a X-ray amorphous phase.

Under the literary data amorphous oxide ZrO₂ crystallizes at temperature 393°C, Y₂O₃ is the effective stabilizer of amorphous condition ZrO₂ and leads to rise in temperature of crystallization of amorphous solid solution Zₐ₀.₉₅Y₂O₁₋₉₇₅ to 407–442°C [21]. However, in work [22] temperature of crystallization amorphous zirconia received by thermal decomposition of a carbonate has made 350°C. In our case there was a decrease in temperature of crystallization concerning known [21] almost on 220°C that unequivocally specifies in essential influence of the dimensional factor on temperature of crystallization X-ray amorphous components as the average size of crystalline particles in it is considerable below (1-2 nm), than at amorphous ZrO₂ in work [22].

We experimentally found for the first time a ferromagnetic condition at room temperature in NPS YSZ [16]. From the data Figure 5 it is visible that magnetization of saturation NPS YSZ makes 0.067 emu/g. Almost at the same time with us, in the review [23] were provided magnetic characteristics of thin films of YSZ to room temperature (3.5 × 10⁻⁷ emu/g), this is on 5 orders lower than size of magnetization of NP YSZ established by us. Recently, one more experimental confirmation of a ferromagnetism of ZrO₂ is given in C. N. R. Rao work [24]. Ferromagnetism is found out in nanoparticles tetragonal phase ZrO₂ with the sizes 5–8 nm at temperatures 673 and 873 K (3.97 × 10⁻³ and 1.93 × 10⁻³ emu/g, accordingly), received by hydrolysis ZrOCl₂ at a microwave irradiation in water solution NaOH.

The experiments with gadolinium-doped ceria confirmed the earlier results and demonstrated the possibility of producing powders with Sₜ of about 190 m²/g. The synthesized powders had a large-specific surface and were agglomerated to complexes 20 to 600 nm in size similar to those of YSZ (Figure 6).

The X-ray diffraction analysis showed that two phases, a mechanical mixture of CeO₂ and Gd₂O₃, were present in the material of the target. Given below are characteristics of their structures.

(i) CeO₂: a cubic lattice of the fluorite type with the spacing a = 5.413 Å; the CSR 84 nm in size; and the concentration of ≈81 weight %;

(ii) Gd₂O₃: a cubic lattice of the Mn₂O₃ type with the spacing a = 10.89 Å; the CSR 124 nm in size; and the concentration of ≈18 weight %.

Remarkably, the value of the lattice spacing of the phase at hand is considerably different from the corresponding values reported in the literature (10.797 Å ÷ 10.813 Å).
According to the analysis, the material is complicated and does not lend itself to an unambiguous interpretation. A model of one phase—a solid solution of gadolinium in CeO$_2$—is considered as a first approximation. Figure 8 shows parameters of the diffraction pattern as refined by the method of full-profile analysis. The following characteristics of the structure were determined: the lattice spacing $a = 5.431\text{Å}$ and the CSR size equal to 3.7 nm. It should be noted however that the estimated spacing corresponded to the Gd$_2$O$_3$ concentration equal to or larger than 20 mole % (this value already corresponds to the two-phase region of the concentrations). Furthermore, the sharp fragments of the diffraction peaks, which are seen in the diffraction pattern, are not described by the Lorentz function, which is typical of dispersed powders. Therefore, a more complex model needs to be considered.

A model of two phases was considered as a first approximation.

(1) A solid solution of cerium in the cubic lattice of Gd$_3$O$_5$, a fine crystalline phase.

(2) A solid solution of gadolinium in the cubic lattice of CeO$_2$, a coarse crystalline phase.

Results of the refinement procedure are given in Figure 7. The following characteristics of the structure were determined.

(i) A solid solution of cerium in the cubic lattice of Gd$_3$O$_5$: the spacing $a = 10.88\text{Å}$; CSR 3 nm in size; the concentration of $\approx 97$ weight %.

(ii) A solid solution of gadolinium in the cubic lattice of CeO$_2$: the spacing $a = 5.415\text{Å}$; CSR $\approx 60$ nm in size; and the concentration of $\approx 3$ weight %.

Thus, the X-ray diffraction analysis demonstrated that the two-phase nanopowder, which was synthesized in the cerium-gadolinium system, can be interpreted as resulting from an incomplete reaction (a mechanical mixture was the initial material). This interpretation does not apply to the zirconium-yttrium system (one phase was present in the initial material). The insynthesis formation of two phases, which differ by the CSR size and the chemical composition, is of principal significance and is connected with the synthesis method used.

Using installation Nanobeam-2 were synthesized new X-ray contrast substance (XRCS), consisting of difficult oxides tantalum and rare-earth elements (LaTaO$_4$ and YTaO$_4$ [25]), and their comparison with the same of materials in a microsize ($S_{BET} \approx 0.04\text{ m}^2/\text{g}$) has been executed. Received NP had $S_{BET} \sim 165–185\text{ m}^2/\text{g}$. According to PEM VR are received poorly agglomerated NP consisting of units, formed crystal LF in the size of $\sim 3–5$ nanometers, with very narrow distribution of particles on the size which in turn form agglomerates in the size to several hundred nanometers. At all NP there are large particles of the micron size. Compared XRCS in 5 wt. % water suspension were placed in glass bottles in capacity of 10 mL and irradiated using the RUM-20 M X-ray device. For dose measurement the X-ray film was used on usual technology. Density of dimness was measured on MD-100 microdensitometer.

It is established that X-rays contrast of images of the bottles containing nanoslips above, than the bottles containing suspensions with micropowders on 20–30%. Thus, it was found that X-rays contrast of XRCS depends not only from chemical, but also granulometric structures of substances [25].

At production nanopowder Al$_2$O$_3$ and Al$_2$O$_3$-Cu(Al) in vacuum was found that with increase in concentration of copper in NPS Al$_2$O$_3$-Cu the size of a specific surface decreases (Table 2). The greatest $S_{BET}$ is received for weak doped NP Al$_2$O$_3$-1,7 wt % Cu, thus its size is close to record indicators for NPS pure Al$_2$O$_3$ (340 m$^2$/g [9], 320 m$^2$/g [26]).

XRD it is established (Table 3) that NPs contain on three crystal phases: α-Al$_2$O$_3$ (corundum), γ-Al$_2$O$_3$ (cubic), and θ-Al$_2$O$_3$ (monoclinic) phases. In NPs, doped Cu, was found out crystal soured copper. At all NPs is present as well the amorphous component. In Table 4 values of the periods of crystal lattices of the revealed phases are resulted.
Figure 8: (a) Chains of particles (NPS Al₂O₃·1.7 wt. % Cu); (b) aggregates of particles about 5 nm in size (NPS Al₂O₃·1.7 wt. % Cu).

Table 2: Specific surface NPS Al₂O₃ and Al₂O₃-Al (Cu).

<table>
<thead>
<tr>
<th>Composition NPs (PEB)</th>
<th>Al₂O₃</th>
<th>Al₂O₃-Al</th>
<th>Al₂O₃-1.7 %Cu</th>
<th>Al₂O₃-11.2 %Cu</th>
<th>Al₂O₃-17.2 Cu</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sₜᵉᵗ, m²/g</td>
<td>269.69</td>
<td>197.80</td>
<td>338.25</td>
<td>80.71</td>
<td>68.71</td>
</tr>
</tbody>
</table>

Table 3: Relative concentrations of the crystalline phases and average values of CSR.

<table>
<thead>
<tr>
<th>Composition NPs (wt % Cu)</th>
<th>α-Al₂O₃ (Corundum)</th>
<th>γ-Al₂O₃ (cubic)</th>
<th>θ-Al₂O₃ (monoclinic)</th>
<th>Cu₂O</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Concentration, wt %</td>
<td>Concentration, wt %</td>
<td>Concentration, wt %</td>
<td>Concentration, wt %</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>65 (3)</td>
<td>78 (2)</td>
<td>24 (3)</td>
<td>21 (2)</td>
</tr>
<tr>
<td>Al₂O₃-Al</td>
<td>64 (3)</td>
<td>68 (2)</td>
<td>25 (3)</td>
<td>26 (2)</td>
</tr>
<tr>
<td>Al₂O₃ (1.72)</td>
<td>28 (3)</td>
<td>75 (3)</td>
<td>45 (3)</td>
<td>32 (2)</td>
</tr>
<tr>
<td>Al₂O₃ (11.19)</td>
<td>60</td>
<td>45</td>
<td>23</td>
<td>28</td>
</tr>
<tr>
<td>Al₂O₃ (17.22)</td>
<td>55</td>
<td>43</td>
<td>26</td>
<td>28</td>
</tr>
</tbody>
</table>

Table 4: Lattice spacing of the phases in Al₂O₃-Al (Cu) NPs.

<table>
<thead>
<tr>
<th>Sample (wt % Cu)</th>
<th>α-Al₂O₃</th>
<th>γ-Al₂O₃</th>
<th>θ-Al₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>a, Å</td>
<td>c, Å</td>
<td>a, Å</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>4,761 (2)*</td>
<td>13,003 (5)</td>
<td>7,918 (4)</td>
</tr>
<tr>
<td>Al₂O₃-Al</td>
<td>4,759 (2)</td>
<td>12,993 (5)</td>
<td>7,905 (4)</td>
</tr>
<tr>
<td>Al₂O₃ (1.72)</td>
<td>4,759 (2)</td>
<td>12,987 (5)</td>
<td>7,909 (4)</td>
</tr>
<tr>
<td>Al₂O₃ (11.19)</td>
<td>4,770 (5)</td>
<td>12,997 (7)</td>
<td>7,914 (6)</td>
</tr>
<tr>
<td>Al₂O₃ (17.22)</td>
<td>4,761 (5)</td>
<td>13,024 (7)</td>
<td>7,930 (4)</td>
</tr>
<tr>
<td>Literary data from file PDF-2</td>
<td>4,7587</td>
<td>12,9929</td>
<td>7,939</td>
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<tr>
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<td>00-046-1212</td>
<td>00-050-0741</td>
<td>00-023-1009</td>
</tr>
</tbody>
</table>

* In brackets is an absolute error of measurement XRD.

Thus the strongest changes was observed for low-temperature metastable γ-phases. On absolute size the periods of a lattice scale of a phase much less than their known values under the literature, the least correspond not doped or doped aluminum to a case. It is not surprising-synthesis passed in vacuum conditions, hence, naturally to expect presence in a material of a considerable quantity of oxygen vacancies.

The general in morphology of samples Al₂O₃-Al(Cu) consists in the following.

1. Large shapeless and spherical particles in the size up to 10 micron, most likely it is target splinters.
2. Shapeless amorphous particles in the size from 40 nm to 1 micron at which the picture of electronic
diffraction of the allocated area represents characteristic halo.

(3) Chains of particles in the size of 50–70 nm, each of which represents agglomerates from particles in the size about 5 nm (Figure 8) with very narrow distribution of particles on the size.

(4) The crystalline particles covered with an amorphous layer in the thickness from 2 to 10 nm (Figure 9).

(5) TEM with the nuclear permission were received only at samples Al₂O₃-Cu, and at samples Al₂O₃-Al; it has not turned out, because of presence on a surface of an amorphous layer.

Room-temperature ferromagnetism has been observed in Cu, Al-doped Al₂O₃. As a whole the level of magnetization Al₂O₃-Cu NPs were in ten times more than the level of magnetization Al₂O₃ NPs fabricated by thermal heating of aluminum hydroxide [27]. Most likely it is connected with presence of magnetic ions Cu²⁺, and also the big defective structure of NPs, formed as a result of evaporation PEB, than at a chemical method [27]. At increase in concentration Cu-specific magnetization increases in structure of NP (Figure 10), however, not proportionally with ferromagnetic impurity (Table 5).

Besides, this data is visible to proportionally ferromagnetic impurity that concentration of copper in NPS is essential more than in initial targets that is connected with the big distinction partial pressure of copper and corundum.

Let us notice that induced by oxygen vacancies ferromagnetism in NPs Al₂O₃ is found recently out in [28]. Amorphous and crystal nanoparticles Al₂O₃ have been received by a zol-method with the subsequent annealing at different temperatures. Magnetic measurements have shown that all nanoparticles showed internal at room-temperature ferromagnetism and magnetization of samples grew $1,25 \times 10^{-3}$ emu/g after annealing in vacuum though bulk sample Al₂O₃ was paramagnetic. Record indicators of magnetization (4.6 emu/cm³) have been established in the thin films Al₂O₃ [29] received by magnetron dispergation on Si-substrate with the subsequent reannealing in vacuum at 800°C. Zheng et al. [29] have explained an origin ferromagnetism formation oxides vacancies on the interface between thin films Al₂O₃ and a silicon substrate. Thus, experimental acknowledgement ferromagnetism in Al₂O₃ in works [27–29] will be coordinated with results of our magnetic measurements in NPS Al₂O₃-Cu [30, 31].

The idea of universality superficial ferromagnetism in the inorganic NPs, put forward in works [24, 27], has found the acknowledgement in our work c nanopowder on the basis of ZnS [32] and further works [33, 34].

Results of measurements of magnetization NPS ZnS and ZnS-Al, evaporated in vacuum by means of PEB [32], are resulted in a Figure 11. Our data will quantitatively be coordinated with results of work [33] where also, experimentally and it is theoretically proved that vacancies Zn can induce internal ferromagnetism (0.05 emu/g) in not doped ZnS nanowires. Magnetization of saturation nanocrystalline
Table 5: Composition of the targets and the elemental composition of Al$_2$O$_3$-Al (Cu) NPs as determined by the ICP method.

<table>
<thead>
<tr>
<th>Target composition (wt % Cu)</th>
<th>Cu</th>
<th>Fe</th>
<th>Cr</th>
<th>Ni</th>
<th>Si</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$_2$O$_3$</td>
<td>&lt;0.02</td>
<td>0.0607 ± 0.01</td>
<td>0.0030 ± 0.01</td>
<td>&lt;0.00 ± 1</td>
<td>0.3857 ± 0.1</td>
</tr>
<tr>
<td>Al$_2$O$_3$-Al</td>
<td>0.0086 ± 0.0008</td>
<td>0.0387 ± 0.004</td>
<td>0.0005 ± 0.0001</td>
<td>0.0005 ± 0.0001</td>
<td>0.46 ± 0.1</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (1,72)</td>
<td>1.72 ± 0.3</td>
<td>0.1335 ± 0.03</td>
<td>0.0018 ± 0.0003</td>
<td>&lt;0.0001</td>
<td>0.5080 ± 0.1</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (11,19)</td>
<td>11.19 ± 1.1</td>
<td>0.0509 ± 0.005</td>
<td>0.0009 ± 0.0001</td>
<td>0.0009 ± 0.0001</td>
<td>0.10 ± 0.3</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (17,22)</td>
<td>17.22 ± 1.7</td>
<td>0.0628 ± 0.006</td>
<td>0.0015 ± 0.0002</td>
<td>0.0008 ± 0.0001</td>
<td>0.31 ± 0.03</td>
</tr>
</tbody>
</table>

Figure 12: Various structures of the wrong form of ZnO and Zn-ZnO.

powders received by hydrothermal synthesis, after annealing on 500°C in steams Zn has made $8 \times 10^{-3}$ emu/g [34] that is close to indicators of magnetization of ours NPS [32]. Ferromagnetism in doped by not magnetic elements (Cu, C, N) ZnS is predicted in theoretical works [35–37] and it is experimentally established in powders ZnS-Cu (MS ~ $7 \times 10^{-3}$ emu/g) [38] and doped by transitive metals (Cr, Mn, Fe, Co, Ni) [39].

At evaporation of targets from ZnO (at residual pressure $\sim 4$ Pa) managed to be received NPS Zn-ZnO with $S_{BET} = 56.7$ m$^2$/g variable structure and various morphology (Figure 12).

Large NPS consist of crystalline particles, differ the big morphological variety, perfection of crystal structure, and essential disorder of the sizes. Separate particles (Figure 13) are core-shell Zn-ZnO structure. It is necessary to notice that in work [40] carbothermic synthesis purposely received hexagonal nanostructure ZnO similar on received by us.

4. Conclusion

The created installation allows to fabricate NPs oxides with high-specific surface at productivity till 12 g/hour and specific expenses of energy $\sim 112$ kW*h/g (nearby 5 energy sublimations). At all powders are present fine-crystalline and the coarse-crystalline fractions differing in the size CSR, and also amorphous component. Formation amorphous components in majority of NPs, received by the methods of electron evaporation with use pulse [31, 41, 42], or continuous beam [43], are their prominent features that makes methods of electronic evaporation very attractive to production and studying of properties NPs in interrelation with their amorphous condition. Formation amorphous components in majority of NPS, the methods of electron evaporation received with use pulse [31, 41, 42], or a continuous beam [43], are their prominent features that makes methods of electronic evaporation very attractive to production and studying of properties NPs in interrelation with their amorphous condition.
Powders have fractal structure, consist of units in the size from tens to several hundreds nm formed by crystal NP in the size about 3–5 nanometers, with very narrow distribution of particles on the size. NPs have high deficiency of structure that was reflected in their magnetic properties.

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**References**


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