Structure and optical properties of nanoparticles obtained by pulsed laser ablation of copper in gases

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Abstract. Nano-powders of different composition and structure were obtained by nanosecond pulsed laser ablation (Nd:YAG laser, 7 ns, 1064 nm, 20 Hz, 200 mJ) of a metallic copper target in a simple flow reactor at atmospheric pressure. The crystal structure, nanoparticles’ shape, and surface morphology were shown to depend on the gaseous media used (Ar, N2, CO2). CuO nanopowders with a specific surface area 50 m^2/g were synthesized by pulsed laser ablation in carbon dioxide and after annealing in air. The optical properties of the obtained powders were also investigated.

1. Introduction
At the present time, changes in the materials’ properties occurring at their transition to the nano-sized state are investigated intensively. Nanostructured materials are solids with sizes in the range of 1-100 nm. They include nanocomposites, aggregated nanoparticles (NPs), clusters, nanocrystalline thin films, (0-4) D materials, and semiconductor nanostructures (quantum dots). Due to the reduced size (quantum-size effect), nanostructured materials have electronic, magnetic and chemical properties different from that of the bulk materials. This is connected with an increase in the surface energy and leads to the increased activity and mobility of NPs. Such materials are promising for use in heterogeneous catalysis [1], gas sensing [2] and solar cells [3] technologies, microelectronics [4], nonlinear optics [5], and biomedicine [6].

Nanomaterials based on copper and its compounds, primarily oxides, are of great interest. Different copper-containing materials are used depending on potential applications. Metallic copper has high conductivity, large extinction cross section, high photosensitivity and low cost. At the same time, copper oxides nanoparticles (CuO and Cu2O) are applied in optics and electronics [7]. Variable valency determines the prospects for copper-containing materials in catalysis. Thus, CuO nanoparticles are more often used in oxidative catalysis [1], and Cu2O nanoparticles are applied in photocatalytic and solar energy systems [8, 9]. Recently, copper NPs have attracted a great interest in biomedicine. Copper-containing nanoparticles were shown to antibacterial activity along with low toxicity [10] and excellent anti-pathogenic properties [11, 12].

There are various methods to produce Cu-containing NPs depending on the range of their size, shape, surface morphology and chemical composition [13]. Among the existing methods, there are chemical processes, such as wet synthesis (from copper hydroxides), sol-gel synthesis, and physical methods, namely thermal sputtering, magnetron sputtering, pyrolysis, etc. However, these methods have a number of drawbacks, such as the use of expensive precursors and complicated equipment. Therefore, the development of green and cost-effective processes for the controlled synthesis of new
nanostructures is of high importance. Pulsed laser ablation (PLA) is a convenient and widely used approach to prepare diverse NPs [14]. Its advantages include technical simplicity and versatility (one-stage process, a wide range of materials used); obtaining of pure nanoparticles that are free of contaminants; and the size control for NPs synthesized.

Pulsed laser ablation is based on a layer-by-layer removal of the material from the surface of the target under powerful short laser pulse in various media. Pulsed laser ablation in liquids (PLAL) has become widespread in the last 20 years as a method of nanoparticle synthesis. During PLA in liquid, the processes of assembly, generation, transformation and condensation for NPs are improved. However, a screening layer is formed consisting of the agglomerated colloids in the long-term ablation. This hinders production of the significant amounts of NPs [14]. Also, to obtain nanopowders via PLA in liquid, additional stages of precipitation and drying of the dispersions are necessary. Therefore, for some applications, it may be appropriate to obtain nanoparticles with PLA in gas or vacuum (PLAG).

It should be noted that since the first lasers appeared, the PLA was widely used for laser processing of materials in vacuum and gas media. Cutting, scribing, modification of the target surface [15], etc. were performing. The laser ablation in gas or vacuum is used for mass spectrometry [16] and the preparation of thin films [17]. Recently, the PLAG started to be used for the synthesis of NPs. For example, the syntheses of NPs by PLAG were done in various gases at low [18, 19] and atmospheric pressure [20, 21, 22]. In the work [23], we carried out a comparative study of nanoparticles obtained in water and air. These studies showed that PLAG can compete with PLAL as a method for nanoparticles synthesis. For example, in a gas, it is possible to better control the chemical reactions of the target material with the medium in order to obtain pure metal particles upon ablation of the active metals [21, 22]. Experimental and theoretical studies of particle formation, plasma dynamics, and the influence of gaseous media on the structure and properties of the resulting NPs are necessary for the development of synthesis methods NPs by PLAG.

To date, we are not aware of studies on NPs obtained by PLAG of metallic copper at atmospheric pressure, although there are many studies on the synthesis of copper-containing particles in liquid phase, for example [24, 25].

The purpose of this work was to prepare NPs by pulsed laser ablation of the bulk copper target in gaseous media at atmospheric pressure and to study their structure and optical properties. We used ordinary atmospheric air and inexpensive and easily available technical gases (CO₂, N₂ and Ar).

2. Experimental Part

2.1. Synthesis of nanomaterials

Nanoparticles were obtained by the pulsed laser ablation method in various gas environments. The scheme of the reactor used is shown in figure 1. The metallic copper target (99.95%) was placed in a glass container, which was filled with gas. Four different gaseous media were used: air, argon, nitrogen and carbon dioxide. The gas flow rate through the reactor was of 10 ml/min to maintain constant concentration of components. A filter was installed at the outlet of the gas system to prevent the release of nanoparticles into the atmosphere. The irradiation of the fundamental harmonic of Nd:YAG laser (LS-2132UTF, LOTIS TII, Belarus) was used (wavelength 1064 nm, pulse duration 7 ns, pulse energy 200 mJ, pulse repetition rate 20 Hz). The beam radiation was focused at the target surface by a long-focus lens with F=500 mm through the optically transparent window of the reactor. The power density of the laser irradiation on the target surface was ~ 0.5 GW/cm². The reactor with the target was moved in the XY plane perpendicular to the optical axis by means of two line translators’ 8MT173-100 (Standart Ltd., Lithuania) for the uniform removal of the target surface. The obtained NPs spontaneously settled on the walls of the reactor and were mechanically collected after 360 min of PLA. Four powder samples were obtained in different gaseous media under equal conditions. The samples were marked according to the gas used: Cu_Air, Cu_Ar, Cu_N₂ and Cu_CO₂.
The obtained nanoparticles were annealed in air at 260 °C for additional oxidation. The required annealing temperature was previously determined based on previous results obtained by differential scanning calorimetry (DSC).

2.2. Composition and structure investigation
The phase structure of the materials was investigated using X-ray diffractometer XRD-6000, Shimadzu (Japan). Diffraction patterns of powders were indexed and the percentage of phases was calculated using the PDF-4 database. The morphology of the surface of the copper-containing nanoparticles was studied using a scanning electron microscope VEGA 3 SBH, Tescan (Czech Republic). Dimensional characteristics of NPs were determined using transmission electron microscopy using a CM 12 microscope, Philips (Netherlands), at the accelerating voltage of 120 kV. For this, the powders were dispersed in ethanol and dropped onto the copper grid coated with carbon thin film. Specific surface area and porosity measurements were realized on an automated sorption system TriStar II (3020), Micromeritics (USA). A volumetric mode of the sorption method was used. The specific surface area was calculated from the isotherm of low-temperature nitrogen vapour sorption.

3. Results and Discussion
The results of XRD analysis of prepared powders are shown in figure 2 and in table 1. It can be seen that PLA of copper in air formed NPs containing 74% of the Rouaite –monoclinic phase of copper trihydroxy nitride Cu₂NO₃(OH)₃ (PDF Card No. 00-045-0594). Also in the Cu_Air sample, a cubic phase of copper (I) oxide (PDF Card No. 01-080-3714) and a monoclinic phase of copper (II) oxide (PDF Card No. 04-007-0518) were found. In addition, a low-intensity peak of the cubic phase of metallic copper is present in the diffraction pattern (PDF Card No. 04-002-8854). When the sample was stored, the particles were found to grow bigger, the crystallinity degree of the Rouaite increased, the cuprous oxide transformed into cupric oxide.

The Cu_Ar powder obtained in argon is represented by NPs consisting of Cu₂O and Cu cubic phases and a monoclinic CuO. About 62% of the cubic phase is formed via PLA of copper in nitrogen. Also, there were phases of Rouaite, copper oxide, and metallic copper in sample Cu_N₂. The powder obtained by PLA in carbon dioxide was characterized by the presence of 89% of Cu₂O cubic phase and 11% of metallic copper. In addition, a low-intensity peak of the monoclinic phase CuO (less than 1%) can be found in the Cu_CO₂ sample.
IR spectra (figure 3) provided additional information on the composition of produced NPs. For samples Cu_Air and Cu_N₂ (containing Rouaite according to XRD) Cu-OH₂ bonds appear in the spectra (wagging, rocking, twisting modes of OH groups in the region around 884, 808 and 677 cm⁻¹, respectively) [26].

**Table 1. Phase composition of prepared powder samples as measured by XRD.**

| Media | Phase, % |
|-------|----------|
|       | Cu      | Cu₂O   | CuO    | Cu₂(NO₃)(OH)₃ |
| CO₂   | 11      | 89     | <1     | –             |
| N₂    | 9       | 62     | 14     | 15            |
| Ar    | 7       | 49     | 44     | –             |
| Air   | <1      | 20     | 6      | 74            |

**Figure 2.** X-ray diffraction patterns for powders obtained by PLA in different gaseous media.

The band at 1631 cm⁻¹ belongs to the in-plane bending of the OH group. The bands with maxima at 1427, 1340 and 1045 cm⁻¹ indicate the presence of a monodentate nitro group bound to metallic copper by a single bond (Cu-ONO₂). Moreover, the bands at 1330, 884 and 597 cm⁻¹ corresponding to bidentate ligand – NO₂, are present in the spectrum of the sample Cu_N₂ powder [26]. For the sample Cu_CO₂ the bands with maxima at 1479, 1402, 1041, 835 and 685 cm⁻¹ confirm the presence of bidentate and/or bridged carbonates on the surface of NPs [27, 28]. The presence of copper (II) oxide in the IR spectra of samples Cu_Ar and Cu_N₂ is confirmed by the bands with maxima at 541, 510 and 480 cm⁻¹ belonging to Cu-O oscillations [29, 30].

**Figure 3.** AT-FTIR spectra of powders generated in different gaseous media. ν – stretching, δ – in-plane bending or deformation, ρₜ – wagging, ρᵣ – rocking, ρₜ – twisting.

It is worth noting that the annealing in the air at 260°C led to the Rouaite phase decomposition, copper (I) oxide and metallic copper oxidation. During heat treatment, all the samples studied were completely converted into copper (II) oxide with a monoclinic structure.
The SEM images of the powders and their specific surface values are shown in figure 4. It can be seen that the initial nanopowders showed non-faceted shape. After annealing, the nanoparticles grew bigger and acquired faceted shape. These changes are better observed in the case of plate-like NPs obtained by PLA in air.

**Figure 4.** The SEM images of the powders obtained by PLA in different gaseous media before and after thermal treatment.

The as-prepared sample Cu_CO₂ exhibited the largest specific surface area of 78 m²/g. After annealing in air at 260°C its S_BET decreased to 51 m²/g. The lowest BET surface area was determined for sample Cu_Air, being 8 m²/g. After annealing the particles of the sample agglomerated, and S_BET decreased to 5.85 m²/g.

Figure 5 shows the TEM images of sample Cu_CO₂ before (a) and after annealing in air at 260°C (b).

**Figure 5.** TEM images of sample Cu_CO₂: before (a) and after annealing in air at 260°C (b).

UV-Vis absorption spectra of the initial and annealed (260°C, 4 h) powders are presented in figures 6 and 7, respectively. The initial powders were characterized by an absorption band in the red region of the spectrum with a maximum at 590-620 nm. This region belongs to the surface plasmon resonance of metallic copper NPs [32]. Note that intensity of this peak increases with increase in the metal phase content (table 1). This may be associated with the presence of oxide phase on the metal surface [32] and with the presence of copper nitrite crystalline hydrates. For the powder obtained in
air, intensive absorption in the near UV region was observed that associated with the copper nitrate absorption [33].

In the shortwave spectral range of 350 and 450 nm, the samples have implicit transitions related to absorption of oxide phases (Cu$_2$O and CuO). According to the literature, cuprous oxide and cupric oxide can absorb in this area (calculation shows that, from the point of view of the band theory, the absorption bands of Cu$_2$O are explained by the formation of exciton pairs, and the absorption bands of cupric oxide are considered from the point of view of field theory ligand) [34-37]. Thus, the absorption bands of both CuO and Cu$_2$O were present in the spectra of all samples. The absorption spectra of all annealed powder samples were found to be similar, which confirms the formation of only the monoclinic CuO phase (figure 7). They have a broad absorption band with a maximum at 350 nm, which is related to the CuO phase [36, 37].

![Figure 6. DRS data for the powders obtained by PLA in different gaseous media.](image1)

![Figure 7. DRS data for the powders after thermal treatment in air (260°C, 4 h).](image2)

To explain the results obtained, we consider possible physicochemical processes during laser ablation of the copper target in various gases. Local heating of the target to a temperature of more than 5000 °C occurs under our experimental conditions (7 ns, 0.5 GW/cm$^2$) according to the equation for thermal diffusivity [38]. Thus, the formation of plasma from the target material and components of the gaseous medium occurs during ablation. We assume that the processes during ablation in various gases proceed as follows: the metal plasma formation → the formation of gas plasma and the gas decomposition → plasma-chemical reactions → relaxation after removing from the reactor.

The air components ionization occurs during PLA in air [39, 40]. Copper clusters react with the air plasma (NxOy) components. The oxidation reaction of copper clusters with activated oxygen of air can also occur. A light green-blue powder was obtained after ablation in the reactor. It consisted mainly of copper nitrates with impurities of oxides and metal particles. The powder becomes dark blue after interaction with air (water vapour), that is, it turns into Rouaite.

The nitrogen molecule ionization [41] happens at ablation in N$_2$. This process is less effective. We assume the formation of copper nitride Cu$_3$N as a result of interaction between Cu clusters and active nitrogen species. This is confirmed by the formation of a gray-green powder in the reactor. Copper nitride passes in Rouaite upon contact with air, and pure and highly active metallic NPs of copper are oxidized by atmospheric oxygen.

The oxidation of Cu NPs during ablation in inert argon occurs after the extraction of the powder from the reactor upon contact with atmospheric oxygen. Reactions with uncontrolled oxygen impurities are also possible.
Ionization and thermal decomposition of CO$_2$ occur with the formation of CO and oxygen. Oxidation of copper clusters occurs. These processes are possible because copper is an effective catalyst [42] which participates in the redox reactions of CO $\leftrightarrow$ CO$_2$.

4. Conclusion
Nanostructures with different composition were synthesized by pulsed laser ablation of a copper target in a flow-through gas reactor containing various gaseous media (air, Ar, N$_2$, CO$_2$). The following results were obtained and the assumptions were made about the ongoing physicochemical processes.

- It has been established that copper effectively interacts with nitrogen, its compounds and water vapour, which leads to the formation of coarse particles of Rouaite – Cu$_3$NO$_3$(OH)$_3$ during PLA in air.
- PLA in a nitrogen environment confirms the possibility of direct interaction between nitrogen and copper. However, this process is not effective in the absence of oxygen. Only about 15% phase of Rouaite forms as a result. NPs interact with oxygen and air-water vapour after being removed from the reactor. We think that Rouaite is formed from copper nitride particles, and copper oxides are formed from metal particles.
- The formation of Rouaite does not occur drying PLA in CO$_2$ and Ar. Oxidation copper takes place directly during ablation in CO$_2$. As a result, highly dispersed particles Cu$_2$O (up to 90%) are formed.
- In air, the main oxidation processes occur after the extraction of pure active particles from the reactor, although some of the particles may be oxidized during the ablation process due to O$_2$ impurity in argon. As a result, particles of copper (I) and (II) oxides form in nearly equal proportions.

After annealing in air at 260°C, all the powders are transformed into a bivalent oxide with a monoclinic structure. The particles obtained by PLA in CO$_2$ have the largest specific surface area.

Thus, ultrafine nanopowders were obtained by PLA in CO$_2$ at atmospheric pressure. They mainly consisted of Cu$_2$O or CuO (during heat treatment at 260°C) These nanopowders will be tested in heterogeneous catalysis and as antibacterial agents in biomedicine.

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