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PARAMETERS OF NANOSECOND OVERVOLTAGE DISCHARGE PLASMA IN A NARROW AIR GAP BETWEEN THE ELECTRODES CONTAINING ELECTRODE MATERIAL VAPOR

Parameters of the nanosecond overvoltage discharge plasma in an air gap of $(1 \div 5) \times 10^{-3} \text{ m}$ between the electrodes, which contains the vapor of an electrode material (Zn, Cu, Fe) injected into plasma due to the ectonic mechanism, have been studied. The dependences of those parameters on the ratio $E/N$ between the electric field strength $E$ and the particle concentration $N$ in the discharge are calculated for the “air–copper vapor” system, by using the numerical simulation method.

Keywords: nanosecond discharge, air, radiation emission by atoms and ions, plasma parameters, zinc, copper, iron.

1. Introduction
Nano- and subnanosecond high-current discharges in the air gap between metal electrodes are a source of ultraviolet (UV) radiation emitted by atoms and ions of an electrode material (copper, iron, molybdenum, tungsten, and others) in a spectral interval of 200–300 nm [1–5]. The vapor of an electrode material is injected into the discharge by microexplosions occurring at the surface of a needle electrode and an electrode on the basis of the sharp edge of a metal foil, as well as inhomogeneities on the surface of metal electrodes. Those microexplosions result in the appearance of ectons [6].

Unlike UV and VUV lamps with the pumping by a barrier or capacitive discharge [7, 8], “point-source” UV lamps on the basis of a vapor of chemical elements can have very small plasma volumes of about 1–5 mm$^3$ at a pulse power of radiation emission larger than 1 W. Therefore, they are applied to the calibration of spectral devices and various optical elements operating in the UV spectral interval. They are also promising in nanotechnology, medicine, and biology. The parameter of those “point-source” lamps have been most completely studied, when exciting an overvoltage discharge 1–2 ns in duration, which is ignited making use of unique pump oscillators [9].

Less studied are the parameters of nanosecond discharges between metal electrodes for systems with a low inhomogeneity of the electric field strength distribution in the discharge gap [10–13]. Such conditions are created, when a system of electrodes of the “sphere-sphere” type is used, and the curvature radius of the working electrode surface is comparable with the distance between the electrodes [14].

The application of bipolar high-voltage pulses allows one to make the device more compact. Furthermore, in this case, the electrode material is spent...
more uniformly, when the electrode is destroyed in a strong electric field by means of microexplosions. As far as we know from the literature, no researches have been carried out till now concerning the emission characteristics of “point-source” UV lamps on the basis of a zinc vapor. When zinc electrodes are used, plasma mainly emits in a narrow spectral interval of 200–215 nm, which is located at the edge of the air transmission interval and is of interest for the application in medicine as a bactericidal radiation source. Since zinc has relatively low melting and evaporation temperatures, a reactor of this type can be promising for synthesizing thin nanostructured zinc oxide films [15]. Researches of such discharges make it possible to design a windowless reactor, which is a source of both the selective UV radiation and nanoparticles of transition metal oxides, including magnetic ones on the basis of iron oxides. Such devices are also promising for application in medicine and biology [16].

In works [17, 18], the results of detailed researches were reported concerning the parameters of the cathode and anode plasma formation in volumes of tens of cubic micrometers at the pre-breakdown phase of a nanosecond overvoltage discharge in the electrode system “tip-plane” filled with air under the atmospheric pressure. The electron concentration in these plasma microformations reaches a value of \((5 \pm 6) \times 10^{23} \text{ m}^{-3}\). After a cathode plasma bunch has been formed and the discharge current in the gap has drastically grown, the cathode plasma bunch unites with the corresponding plasma formation, which moves from the anode. The relation between the emission parameters of plasma and the electrode materials were not quoted in the cited works.

The repetitive pulsed discharge in xenon with a microsecond pulse duration is an effective source of continuous radiation emission in a spectral interval of 120–500 nm (with a possible increase of the lamp efficiency to 20–25% at the pressure \(p(\text{Xe}) = 36 \text{ kPa}\) [19]. By increasing the xenon pressure to 1000 kPa and exciting a nanosecond microdischarge in plasma, a diffuse discharge was obtained in the gap between tips, which emitted in a spectral interval of 200–800 nm [20].

The results of studies of the high-current subnanosecond discharge in gases, generation mechanisms of runaway electrons and accompanying X-rays, as well as the parameters of this plasma and the main applications of such discharges, were summarized in works [21–24]. The conditions and mechanisms of formation of homogeneous plasma aggregates with a high density and a large volume in strongly non-uniform electric fields of the high-pressure multielectrode corona discharge were considered in [25]. The main role under those conditions was played by nonlinear processes in the plasma of heavy inert gases and the effect of strong electron sticking to electronegative halogen-based molecules.

One of the promising applications of nano- and subnanosecond discharges in air can be the synthesis of nanostructures of transition metal oxides (CuO, Cu2O, ZnO, Fe2O3, Fe3O4, and so forth). Nanostructures of transition metal oxides are synthesized, by using methods that are based on the deposition of nanostructures from the gas phase by thermal, laser, or magnetron sputtering. Laser methods for the nanostructure synthesis are characterized by the low yield of a final product, whereas other gas-discharge methods require the application of a vacuum equipment. The capabilities of the synthesis of transition metal oxide nanostructures in the nanosecond overvoltage discharge with the ectonic mechanism of electrode material vapor injection into a material and the nanostructure parameters have not been studied earlier. Therefore, a task to study the conditions for the synthesis of such nanostructures in atmospheric-pressure air, which would be possible making no use of expensive laser and vacuum facilities, is challenging.

In this work, the results of our research concerning the spatial, electrical, and optical characteristics of a nanosecond overvoltage discharge between the electrodes made of transition metals (zinc, copper, iron) in air and under conditions of a rather uniform electric field strength distribution in the “sphere-sphere” discharge gap are reported.

2. Experimental Method and Equipment

The design of a discharge chamber with a system of electrodes, between which a nanosecond discharge was ignited in air, is shown in Fig. 1. The electrodes were fabricated from zinc, copper, or stainless steel and mounted in a discharge chamber 0.03 m³ in volume fabricated from an insulator. The distance between the electrodes was 0.001–0.005 m, so that the discharge was overvoltaged. The air pressure in the
chamber was varied in an interval of 6–303 kPa. The discharge chamber was evacuated to a residual pressure of 10 Pa, but most of the experiments were performed at an air pressure of 101 kPa (1 atm) and in the absence of the output quartz window in a discharge chamber [26, 27].

In order to reduce the electromagnetic influence on the registration system of high-current nanosecond discharge parameters, the experimental chamber with the electrode system was screened with the help of a metal mesh. The diameter of cylindrical metal electrodes was equal to 0.005 m, and the curvature radius of the working electrode tips was 0.003 m. Provided this relationship between the electrode curvature radius and the distance between the electrodes, the electric field strength in the gap was relatively uniform [14]. In order to localize the discharge in the gap between metal electrodes, the latter were inserted into fluoroplastic tubes, so that the length of their non-isolated sections did not exceed 0.003 m. When the electrodes of a nanosecond discharge were used together with the dielectric tubes, an expansion of the discharge glow region was observed [28].

In order to ignite a high-current nanosecond discharge, high-voltage bipolar pulses 50–100 ns in duration and ±(20÷40) kV in amplitude were supplied to one of the electrodes. As a result, a diffuse discharge with a peak current of 50–170 A was ignited between the ends of spherical metal electrodes. The plasma volume did not exceed $5 \times 10^{-9}$ m$^3$. If the distance between the electrodes was $d = 0.001$ m and the air pressure was equal to 1 atm, the discharge gap was overvoltaged, because the electric field strength for an non-overvoltaged gap equals 0.30 kV/m at the air pressure $p = 101$ kPa [29, 30], whereas this parameter reached 3–5 kV/m in our experiments. The pulse repetition frequency was varied in the interval $f = 40÷1000$ Hz.

The voltage pulses across the discharge gap and the discharge current were measured using a broadband capacitive voltage divider, a Rogowski belt, and a broadband oscilloscope 6 LOR-04. The time resolution of this registration system was 2–3 ns. The spatial characteristics of a nanosecond discharge were studied using a digital photo camera. The discharge radiation spectra were registered making use of a monochromator MDR-2, a photomultiplier FEP-106, a DC amplifier, and an electronic potentiometer. The plasma emission was analyzed in a spectral interval of 200–650 nm. The corresponding registration system was calibrated, by using the radiation from a deuterium lamp in a spectral interval of 200–400 nm and from a band lamp in an interval of 400–650 nm.

The radiation emission power of the point-source lamp was measured with the help of an ultraviolet radiometer TKA-PKM, which was tuned to measure in a spectral interval of 200–280 nm. The radiation selection geometry was not taken into account in this case. Therefore, only the average radiation power in a given spectral interval was determined in relative units.

3. Discharge Parameters

Figure 2 demonstrates the registration results for the spatial characteristics of a discharge. From this figure, it follows that a spherical diffuse discharge takes place in the “sphere-sphere” system of electrodes provided the atmospheric air pressure and the distance between the electrodes to be comparable with the curvature radius of the metal electrode. The volume of this discharge strongly depends on the repetition frequency of voltage pulses. The “point-source lamp” operation mode is achieved only at the repetition fre-
Fig. 2. Photos of a nanosecond overvoltage discharge between the copper electrodes at pulse repetition frequencies of 40, 150, and 1000 Hz (from left to right)

frequencies of voltage pulses within an interval of 40–150 Hz.

When the repetition frequency of voltage pulses was increased up to 1000 Hz, the plasmoid diameter grew by 3 to 4 times and covered new areas on the working electrode surface. It can be a result of the residual phenomena in plasma, when the concentration of charged particles has no time to relax to its original level within the time interval between pulses. The spherical shape of the plasma glow region can be associated with the application of dielectric tubes on the electrodes and the accumulation of slow-moving positive ions on the insulator surface, which are formed as a result of the ionization processes near the electrodes and change the configuration of the electric field strength in the interelectrode gap [28]. The second cause for this phenomenon can be the specific features of the streamer formation in strongly overvoltaged gaps of a nanosecond discharge in air. The streamers are generated in the direction perpendicular to the interelectrode gap and the electric field strength vector [9].

The diffuse character of the examined discharge at the atmospheric air pressure is confirmed by the results of researches of the spatial parameters of the transverse nanosecond discharge making no use of a special preionization system, when the discharge was ignited in nitrogen at the pressures $p = 4 \div 101$ kPa taking advantage of a high-voltage modulator with the voltage pulse amplitude $U < 35$ kV [31]. The photos of this discharge together with the registration data of the discharge spatial characteristics making use of a CCD camera showed that the diffuse form of such discharges is retained during the whole voltage pulse at various repetition frequencies. The experimental results [31] testify that a beam of runaway electrons is formed in the strongly overvoltaged nanosecond discharge in nitrogen, only if the nitrogen pressure does not exceed 13 kPa. Therefore, under the available experimental conditions, the main factor determining the diffuse character of the discharge is the X-ray radiation emitted by plasma.

The diffuse character of a plasmoid and its shape close to spherical were similar to those in the photos of nanosecond discharges in air at the distance between the electrodes $d < 0.005$ m and their curvature radius of 0.006 m [29]. The spherical form of the plasmoid can result from the generation of accompanying characteristic and bremsstrahlung X-ray radiation in the strongly overvoltaged discharge gap [30, 32–34], which isotropically illuminates the entire gap. According to data of recently published work [31], a beam of runaway electrons in atmospheric-pressure air and under experimental conditions close to ours can be formed, only if the air pressure does not exceed 13.3 kPa. Therefore, the main factors governing the uniformity of the nanosecond overvoltage discharge, which was studied by us, were the characteristic radiation (the X-ray spectral lines emitted by the metal atoms and the nitrogen atom at the excitation of their inner shells) and the bremsstrahlung X-ray radiation from the anode.

Because of the mismatch between the output resistance of a high-voltage modulator and the discharge impedance, as well as the presence of pulses reflected from the discharge gap with plasma, a voltage pulse consisted of a few small peaks each about 5–10 ns in duration. The duration of the main part of a voltage pulse train reached 50–100 ns. Small bipolar peaks of the voltage pulse had the amplitudes of the positive and negative components up to 15–20 kV. The current pulses of the nanosecond discharge consisted of
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Fig. 3. Pulse power of a nanosecond overvoltage discharge in air between copper electrodes ($p = 101$ kPa, $f = 100$ Hz).

Fig. 4. Plasma radiation emission spectrum of a nanosecond overvoltage discharge in air at $p = 101$ kPa. The interelectrode distance $d = 0.001$ m.

Fig. 5. Plasma radiation emission spectrum of a discharge in the mixture air ($p(\text{air}) = 101$ kPa) + water vapor ($p(\text{H}_2\text{O}) = 130$ Pa). The interelectrode distance $d = 0.005$ m.

A sequence of bipolar current pulses with the amplitudes of the positive and negative components equal to 120–150 A. The total duration of the sequence of current pulses with an amplitude decreasing in time reached 150–200 ns. Such a character of the pulsed energy contribution, which occurs at the mismatch between the output resistance of a high-voltage generator of nanosecond pulses and the discharge plasma impedance, favors the effective sputtering of the electrode material.

The spatial, electrical, and some emission parameters of the bipolar nanosecond discharge between metal electrodes (Zn, Cu, and Fe at $d = 0.5–4$ mm) in the nitrogen atmosphere at a pressure of 1 atm were studied in works [12, 13]. The results obtained there correlate with the results of our experiments.

By multiplying the corresponding data taken from the pulse current oscillogram and the voltage oscillogram, the time distribution of the pulse energy contribution to the nanosecond discharge plasma was calculated. The maximum of the discharge pulse power was observed at the initial stage of the discharge gap breakdown and reached a value of 4 MW (Fig. 3). The integration of the pulse power over the time allowed us to determine the electric energy that was introduced into the discharge plasma during a single sequence of voltage and current pulses. Under the ignition conditions of a nanosecond discharge at an air pressure of 101 kPa and the interelectrode gap width $d = 0.001$ m, the energy introduced into plasma was equal to 105 mJ (Fig. 3). When the electrodes made of zinc or a stainless steel were used instead of copper electrodes, the electrical characteristics of a discharge in air at the atmospheric pressure did not change considerably.

The research of the spectral characteristics of radiation emission by plasma of a high-current nanosecond discharge in atmospheric-pressure air between the electrodes fabricated from transition metals (Zn, Cu, Fe) showed that about 80–90% of the emitted power in a spectral interval of 200–650 nm were concentrated in a spectral interval of 200–220 nm for Zn electrodes, 200–230 nm for Cu electrodes, and 200–260 nm for Fe electrodes. When identifying the spectral lines in the radiation spectra of discharges, reference books [35, 36] were used.

In Figs. 4 and 5, the emission spectra of the nanosecond overvoltage discharge in air and in a mixture of air with a small admixture of water vapor (water vapor was added to obtain radiation spectra with the diagnostic spectral lines of a hydrogen atom), respectively, obtained at various distances between zinc electrodes. At a significant overvolt-
Parameters of Nanosecond Overvoltage Discharge Plasma

age across the discharge gap, when the distance between the electrodes is minimum \(d = 0.001\) m, and in atmospheric-pressure air, two groups of intensive spectral lines emitted by the zinc atom and the zinc ion in a spectral interval of 200–280 nm dominate in the plasma radiation spectra. The most intense spectral lines in group Zn II were the ionic spectral lines at 202.6 and 206.2 nm. In group Zn I, these were the atomic spectral lines at 250.2 and 255.8 nm. Those data correlated well with the spectral parameters of a lamp \([37]\). In comparison with the data of work \([37]\), the UV radiation spectrum in those intervals included much more intense spectral lines of zinc.

As the distance between the electrodes was increased to \(d = 0.005\) m (whereas the air pressure remained constant), the electric field strength in the discharge gap decreased, which led to a decrease of the electron temperature and worsened the conditions of X-ray preionization. The intensity of the spectral lines emitted by zinc atoms and ions in a spectral interval of 200–215 nm practically vanished, and the intensity of the atomic spectral lines of zinc in an interval of 250–280 nm became several times lower. For the plasma of air with small admixtures of zinc vapor and at \(d = 0.005\) m, the spectral lines of zinc in wavelength intervals of 250–280, 330–350, and 460–510 nm became the most intense. If a small admixture of water vapor \((p = 130\) Pa\) was added to air at the atmospheric pressure (see Fig. 6), the radiation spectrum of the discharge demonstrated the low-intensity band of a hydroxyl radical OH at 309.3 nm and the spectral lines of the hydrogen atom in the visible spectral interval.

In the radiation spectrum of the plasma of a mixture of air, water vapor, and zinc vapor, a wide band was registered in a spectral interval of 370–470 nm. This band is not observed in the radiation spectrum of plasma with helium instead of air. The maximum of this band is located in a spectral interval of 410–415 nm. This band can belong to molecules or small clusters on the basis of zinc oxides, which are excited in the gas phase. Those broad bands correlate well with the luminescence bands of nanostructures on the basis of zinc oxide \([38]\).

Figure 6 exhibits the radiation spectra of nanosecond discharge plasma in air at the pressures \(p = 101\) and 303 kPa between copper electrodes in the spectral interval \(\lambda = 200\text{–}230\) nm. As was in the case of less powerful unipolar nanosecond discharges between copper electrodes \([3]\), the spectral lines of copper ions dominate. The obtained spectrum of UV radiation also correlates well with the radiation spectrum of the unipolar subnanosecond discharge with a beam of runaway electrons at a distance of 0.0005 m between copper electrodes \([2]\). In the radiation spectra, the most intense are lines corresponding to transitions from the energy levels of singly charged copper ions \((8.23–9.12\) eV) occurring at the change of electron configuration \(4p–4s\), as well as the spectral lines of copper atoms, which correspond to transitions from highly excited states of a copper ion with energies in an interval of 5.68–7.02 eV.

Figure 7 illustrates a section of the radiation spectrum of nanosecond discharge plasma between the stainless steel electrodes that contains the most intense spectral lines. In Table, the identification of the most intensive spectral lines of iron in a spectral interval of 200–300 nm is presented. The deciphering of the spectrum showed that, in the case of a discharge between stainless steel electrodes, the spectral lines of iron atoms and singly charged iron ions dominate, with the spectral line 249.6 nm FeI being the most intense. Our results correlate well with the corresponding emission spectra of a high-voltage subnanosecond discharge in atmospheric-pressure air registered in the “needle-plane” electrode geometry at \(d = 0.012\) m, when runaway electrons play a significant role \([39]\).
Fig. 7. Emission spectrum of the nanosecond discharge plasma between stainless steel electrodes. The air pressure equals 101 kPa, the interelectrode distance \(d = 0.003\) m, the pulse repetition frequency \(f = 40\) Hz.

In the radiation spectrum of a discharge between a copper and a stainless steel electrode, as well as in the case of a discharge between two copper electrodes, the radiation intensity was maximum for the transitions of singly charged copper ions in a spectral interval of 210–220 nm, and the spectral lines of iron only complemented this spectrum in an interval of 250–300 nm. The most intense of the iron spectral lines was the line 271 nm \(\text{FeI}\). This distribution of the discharge radiation emission between the copper and stainless steel electrodes is mainly associated with different thermophysical properties of indicated metals.

The emission of iron spectral lines in wavelength intervals of 360–400 and 700–730 nm by plasma of a nanosecond discharge in high-pressure helium \((p = 101 \div 505\) kPa) between stainless electrodes \((d = 0.012\) m) at an overvoltage of 100–300% was observed in work [40]. The spectral interval corresponding to shorter wavelengths was not examined in the cited work. Iron atoms were found to emit the most intensively in the high-current diffuse discharge regime, when bright cathode spots were observed on the cathode surface. In air, the high-current diffuse discharge is not ignited.

The dependence of the average power of the UV radiation emission by a nanosecond discharge in a spectral interval of 200–280 nm on the voltage pulse repetition frequency was nonlinear. Its maximum growth was observed in a pulse repetition frequency interval of 40–350 Hz. As this parameter was increased from 35 to 1000 Hz, the average power of the UV radiation emission by the discharge grew by about an order of magnitude. At the same time, the increase of the charging voltage of the working condenser from 13 to 20 kV (at \(f = 35\) Hz) gave rise to the increase of the UV radiation power emitted by a discharge by only a factor of two [5]. The obtained results testify that, in order to increase the average power of the UV radiation emitted by a discharge, the most promising way consists in increasing the voltage pulse repetition frequency.

Under the conditions of described experiments, the ectonic mechanism (the explosion of micropoints on the electrode surface) dominates in the injection of metal vapor into plasma. The corresponding electron concentration in plasma can reach a value of \(10^{16}–10^{17}\) cm\(^{-3}\) [24]. Therefore, the mechanism of appearance of excited metal ions can be governed by the excitation processes of metal ions in the ground state by electrons, and by electron-ion recombination processes. The corresponding effective cross-sections for copper and iron ions are not known at present. The corresponding effective cross-sections for zinc ions have been studied, and they were found to be quite large, namely, \(10^{-16}\) cm\(^2\) [41]. Therefore, it is quite probable that excited atoms and ions of transition metals are formed in the plasma of a nanosecond overvoltage discharge by exciting the singly charged ions of those metals in the ground-energy state by electrons and owing to the recombination process of the doubly charged ions of transition metals by discharge electrons.

The plasma that is dealt with in this work is a complicated object for theoretical studies and even for identification of the spectrum emitted by the nanosecond discharge plasma in air between the stainless steel electrodes \((p = 101\) kPa, \(d = 0.003\) m, and \(f = 40\) Hz)

<table>
<thead>
<tr>
<th>(\lambda, \text{nm} )</th>
<th>Object</th>
<th>(I, \text{rel. un.} )</th>
<th>Lower level</th>
<th>Upper level</th>
</tr>
</thead>
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<td>(a^5D)</td>
<td>(v^5P^0)</td>
</tr>
<tr>
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<td>Fe II</td>
<td>300</td>
<td>(\alpha^6)</td>
<td>(e^6F)</td>
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numerical evaluations, because it contains complex clusters of transition metal oxides and solid micro- and nanoparticles. Thin nanostructured copper oxide films were synthesized on a glass substrate mounted near the discharge. Some spatial and optical parameters of those nanostructures were reported in our work [42].

4. Plasma Parameters

The simulation of plasma parameters was carried out for a nanosecond discharge in atmospheric-pressure air between copper electrodes. Copper vapor was injected into plasma due to the electrode erosion. As was done in work [41], where the plasma in atmospheric-pressure air with small admixtures of copper vapor was investigated and where the partial pressure of copper vapor was estimated to equal 0.1%, the partial pressure of copper vapor for our estimates was chosen to equal 100 Pa. Under the given experimental conditions, the maximum value of the $E/P$ parameter reached a value of about 530 V/(cm torr), which is less than the critical $E/P$ value for nitrogen according to the local criterion of electron escape, 590 V/(cm torr) [30]. Therefore, when simulating the parameters of plasma in atmospheric-pressure air with small admixtures of copper vapor, the standard soft code for solving the Boltzmann kinetic equation to determine the electron energy distribution function [43] was chosen. Unlike similar codes of the BOLCING and BOLCING$^+$ types, which allow the simulation to be carried out only for three chemical elements, the code used by us makes it possible to simulate mixtures of five different components. As a result, the influence of the real air composition on the plasma parameters can be taken into account more completely.

On the basis of the calculated electron energy distribution function, the average energy and mobility of electrons, the specific losses of the discharge power in various electronic processes, and the rate constants of elastic and inelastic electron scattering by copper and argon atoms, and carbon dioxide, oxygen, and nitrogen molecules, as well as their dependences on the reduced electric field (the ratio between the electric field strength $E$ and the total concentration $N$ of copper vapor and gases entering the atmosphere) were determined. The ratio between the indicated components in atmospheric-pressure air was equal to $0.8 : 7.0984 : 0.266 : 159.197 : 593.438$, respectively. The parameter $E/N$ was varied within an interval of 1–1300 Td, which included the experimental values. The set of cross-sections for electronic processes, which were made allowance for in the simulation, and the basic expressions for calculated plasma parameters are given in the code manual [44].

The average energy of electrons in the discharge increased almost linearly from 0.15 to 29.1 eV, as the parameter $E/N$ grew from 1 to 1300 Td. In the experimental interval of the reduced electric field strength 615–820 Td, in which the electrical and optical characteristics of the discharge were studied, the average electron energy varied from 12.5 to 16.0 eV. The electron temperature increased from 145000 to 185600 K, respectively.

According to the results of numerical simulations, the product of the electron mobility and the total concentration of atoms and molecules in the gas mixture changes from $0.7458 \times 10^{24} \text{N} \text{m}^{-1} \text{V}^{-1} \text{s}^{-1}$ to $0.6893 \times 10^{24} \text{N} \text{m}^{-1} \text{V}^{-1} \text{s}^{-1}$ (in m$^{-1}$V$^{-1}$s$^{-1}$ units), when the parameter $E/N$ is varied in an interval of 615–820 Td, which leads to the change of the electron drift velocity from $4.58 \times 10^{5}$ to $5.65 \times 10^{5}$ m/s, provided an electric field strength of $(15.0 \div 20.0) \times 10^{6}$ V/m in plasma. The corresponding values of the electron concentration fall within an interval of $(10.4 \div 8.5) \times 10^{21}$ m$^{-3}$ at a current density of $765 \times 10^{6}$ A/m at the electrode surface $(0.196 \times 10^{-6}$ m$^2$). The calculated electron concentration correlated well with the maximum of the experimentally determined electron concentration in an arc discharge between different composite electrodes, including copper-based ones ($N_\text{e} = 10^{22}$ m$^{-3}$) [45].

The maximum specific losses of the discharge power for inelastic electron processes in the discharge in a mixture of copper vapor with air were observed for a nitrogen molecule and reached 32.7% in the case of excitation of the set of singlet states of a nitrogen molecule at a reduced electric field strength of 820 Td. For the copper atom, the specific power losses of the discharge reached a maximum value of 2.9% at the excitation of its resonant state $^2\text{P}_{3/2,1/2}$ at $E/N = 135$ Td. At a reduced electric field strength of 615 Td, they amounted to 1%, and, at a reduced field strength of 820 Td, they decreased to 0.8%. With the increase of the parameter $E/N$ to 1300 Td, the specific power losses in the discharge reached a maximum
value of 40.2% for the process of nitrogen molecule ionization by electrons.

Figure 8 demonstrates the results of numerical calculations for the dependences of the rate constants of electron collisions with copper atoms on the $E/N$ parameter in a discharge in the mixture of copper vapor with the air $[p(Cu) = 100 \text{ Pa}, p(\text{air}) = 101 \text{ kPa}]$. The rate constants are characterized by large values, $k \approx 10^{-17} - 10^{-12} \text{ m}^3/\text{s}$. This fact is associated with large effective cross-sections of corresponding processes. In an interval of $615-820 \text{ Td}$ for the reduced electric field strength, the excitation rate constants for the $^2P_{3/2, 1/2}$, $^2D_{3/2}$, $^2D_{5/2}$, and $^4F_{7/2}$ states of a copper atom and its ionization are within intervals of $(7.4 \pm 0.6) \times 10^{-13}$, $(0.29 \pm 0.28) \times 10^{-13}$, $(0.23 \pm 0.22) \times 10^{-13}$, and $(0.34 \pm 0.46) \times 10^{-13} \text{ m}^3/\text{s}$, respectively.

The results of numerical calculations carried out for the dependence of the rate constants for electron collisions with nitrogen molecules in a discharge in the mixture of air and copper vapor are shown in Fig. 9. In an interval of $615-820 \text{ Td}$ for the reduced electric field strength, the excitation rate constants for the electronic states of a nitrogen molecule were found to be within intervals of $5.5 \times 10^{-17} - 0.86 \times 10^{-14} \text{ m}^3/\text{s}$ (at $E/N = 615 \text{ Td}$, the states corresponding to curves $1-14$) and $6.4 \times 10^{-17} - 1.5 \times 10^{-14} \text{ m}^3/\text{s}$ (at $E/N = 820 \text{ Td}$, the states corresponding to curves $1-14$). A maximum value of $1.5 \times 10^{-14} \text{ m}^3/\text{s}$ for the excitation rate constant was observed at a reduced electric field strength of $820 \text{ Td}$ in the case of excitation of a set of singlet-singlet states of a nitrogen molecule with an energy threshold of $13 \text{ eV}$.

Large values of excitation constants for the lower electron states of a nitrogen molecule confirm their significant role in the population of the upper energy levels of a copper ion that participate in the formation of the radiation emission by copper plasma in a spectral interval of $200-250 \text{ nm}$. For instance, as was found in work [46], a long-term (for about $2 \text{ ms}$) luminescence of copper atoms was observed in a subnanosecond discharge in a mixture of nitrogen and air at the atmospheric pressure between copper electrodes, which exceeded the discharge current duration by three orders of magnitude. The long-term glow of copper atoms can be associated with the resonance energy transfer from the metastable state $A^3 \Sigma_u^+$ of a nitrogen molecule to the $3d^{10}4d$ level of a copper atom. As a result, a quasimetastable state is formed in a copper atom, through which an effective step ionization of copper atoms with the formation of $\text{Cu}^+$ and $\text{Cu}^{++}$ ions in the ground state can take place. This fact is important for the population of up-
per energy levels of Cu ions at the electron excitation of Cu\(^{+}(0)\) ions and at the recombination of Cu\(^{++}(0)\) ions with slow plasma electrons.

5. Conclusions

To summarize, it is found that a nanosecond overvoltage discharge in the air gap between the zinc or copper electrodes is a selective UV source in spectral intervals of 200–215 and 200–230 nm, respectively. The radiation emission of a discharge between the stainless steel electrodes occupies a relatively wide spectral interval of 200–300 nm. The maximum power contribution to plasma in a pulse was 4 MW, and the energy injected into a discharge during one pulse reached a value of 0.1 J.

The most probable excitation mechanisms of UV radiation emission by atoms and ions of transition metals are the ectonic mechanism of surface erosion of metal electrodes, excitation and ionization of atoms and ions of transition metals by the electron impact, and the formation of excited metal atoms and ions in the processes of electron-ion recombination of singly and doubly charged metal ions with electrons.

The calculation of plasma parameters by numerically solving the Boltzmann equation for the electron energy distribution function showed that, in the experimental interval of reduced electric field strengths (615–820 Td), the average electron energy varies in an interval of 12.5–16.0 eV, and the electron concentration value in an interval of \((8.5\times10^{21}) \times \text{m}^3\). The excitation rate constants for the \(2\text{P}_{3/2,1/2}, 2\text{D}_{3/2,2\text{D}}_{5/2}\), and \(4\text{F}_{7/2}\) states of copper atoms and the constants of their direct ionization by the electron impact are in intervals of \((7.4\times8.6) \times 10^{-13}, (0.29\times0.28) \times 10^{-13}, (0.23\times0.22) \times 10^{-13}, (0.20\times0.23) \times 10^{-15}\), and \((0.34\times0.45) \times 10^{-13} \text{m}^3/\text{s}\), respectively. At \(E/N = 820 \text{ Td}\), the maximum excitation rate constant for a nitrogen molecule, \(1.5 \times 10^{-14} \text{m}^3/\text{s}\), was obtained for the set of singlet-singlet states. The specific losses of the discharge power in inelastic electron collisions with the mixture components were maximum for a nitrogen molecule; namely, 32.7% at the excitation of the set of its singlet states at \(E/N = 820 \text{ Td}\).


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ХАРАКТЕРИСТИКИ І ПАРАМЕТРИ ПЛАЗМИ ПЕРЕНАПРУЖЕНОГО НАНОСЕКУНДНОГО РОЗРЯДУ В ПОВІТРІ З МАЛІМИ ДОМІШКАМИ ПАРИВ ПЕРЕХІДНИХ МЕТАЛІВ

Р е з ю м е

Досліджено характеристики перенапруженого наносекундного розряду в повітрі коли віддаль між електродами складає (1–5)·10^{-3} м та при умовах внесення парів матеріалу електродів (Zn, Cu, Fe) в плазму за рахунок ектоносного механізму. Шляхом числового моделювання розряду в системі “повітря–пари міді” встановлено параметри плазми розряду в залежності від величини відношення E/N, де E – напруженість електричного поля, N – концентрація частинок в розряді.