Membrane Structural Condition and Functional Activity of Peritoneal Macrophages after Gas Discharge Exposure

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The aim of the investigation was to study in experiment the impact of gas discharge (spark plasma and UV radiation of a quartz lamp) radiation on a membrane structural condition and functional activity of peritoneal macrophages.

Materials and Methods. The target of the research was peritoneal macrophages of Wistar rats. A gas discharge Pilimin device series IR-10 and an DBK-9 UV lamp as were used as an operative factor. The suspension of peritoneal macrophages was treated within 30, 60, 300, 600, and 1,200 s. Lipid composition was studied by thin-layer chromatography. Microviscosity in lipid-lipid and protein-lipid interaction areas was analyzed by pyrene fluorescence. Hydrophobicity of membrane hydrocarbon layer was determined by 1,6-diphenyl-1,3,5-hexatrien fluorescence. To assess the functional state of peritoneal macrophages we studied latex particle absorption phagocytosis activity, oxygen-dependent metabolism condition by nitro blue tetrazolium reduction test and by measuring luminal-dependent chemiluminescence.

Results. The study showed phospholipid oxidation to be less intensive under spark plasma radiation than when exposed to UV lamp radiation. Membrane microviscosity in lipid-lipid and protein-lipid interaction areas increases to a greater extent after UV lamp radiation exposure. Spark plasma radiation causes the expansion in the number of cells participating in phagocytosis, the increase of absorbing capacity and oxygen-dependent metabolism of peritoneal macrophages. “Oxygen explosion” time of macrophages decreases after plasma radiation exposure, but increases after UV lamp radiation exposure.

Conclusion. Spark plasma radiation compared to UV lamp radiation contributes to the enhancement of phagocytic activity of peritoneal macrophages in less continuous modes, the effects under study being more pronounced. The study findings enable to reveal optimal modes for functional cell activity after gas discharge exposure.

Key words: gas-discharge devices; peritoneal macrophages; study of spark plasma; UV radiation; membrane microviscosity; membrane hydrophobicity.

The cells of mononuclear phagocytes are of great importance for maintaining body resistance. The immune response intensity and body homeostasis depend on their functional activity [1]. Macrophage structural organization and membrane integrity have an impact on intracellular processes and various metabolic reactions. The condition of cell membrane structural components, their position and interaction determine the biophysical properties of membranes, and therefore, the capabilities of receptor apparatus, integrity and adequate functioning of mononuclear phagocytes [2, 3]. All the above determines the study of physicochemical factors, which enable to enhance the phagocytic activity in infectious or neoplastic processes. In the last decade, biological effects of cold gas discharge plasma have been investigated [4, 5]. There have been found a bactericidal, cytotoxic effect, as well as the activation of mononuclear cells after gas discharge plasma radiation [5–8]. Gas discharge plasma radiation was shown to influence selectively structural-functional components of cell membranes, for instance, it results in no accumulation of molecular products of lipid peroxidation, however, causing oxidative protein modification [9, 10]. When generating gas discharge plasma radiation, UV range, in particular, radical products are formed, which participate in reduction-oxidation reactions. Oxidizers and deoxidizers influence the modification of macromolecules and metabolic processes in a cell [11, 12]. The study and analysis of the effect of gas discharge plasma radiation and UV lamp radiation on macrophage membrane appear to be interesting, it will enable to reveal the mechanism of their modification after the exposure.

The aim of the investigation was to study the impact of gas discharge radiation and UV radiation of a quartz lamp on a membrane structural condition and functional activity of peritoneal macrophages.

Materials and Methods. In vitro experiments were carried out. Peritoneal macrophages of Wistar rats were the research objects. Macrophage suspension...
was obtained 20 min after intraperitoneal administration of sterile saline solution. UV radiation was generated by a DBK-9 UV quartz lamp (Solnyshko, Russia). The spectrum pattern of DBK-9 is monochromatic, the maximum being at λ=254 nm, continuous power is 9 W, an average photon flux is 5.4·10^10 mol(cm^2·s)^{-1}. Spark plasma radiation was generated by an experimental device Pilimin series IR-10 (Skobeltsyn Institute of Nuclear Physics, Lomonosov Moscow State University, Russia) with the pre-set parameters: pulse time is 100 µs, supply voltage is 11 kV, pulse capacitor capacity is 3.3 nF, pulse energy is 5.9·10^{-3} J, frequency is 10 Hz, continuous radiation spectrum with maximum λ=220 nm, average photon flux is (1.26±0.2)·10^{-10} mol(cm^2·s)^{-1}. The suspension of peritoneal macrophages was treated within 30, 60, 300, 600, and 1,200 s. Intact cells served as controls.

At the first research stage we assessed the structural condition of peritoneal macrophage membranes before and after gas discharge exposure: a lipid composition, microviscosity, and hydrophobicity. The lipid composition was analyzed using thin-layer chromatography by successive application of two solvent systems [12]. The chromatograms taken were analyzed and quantitatively processed using GelAnalyzer 2010a. The amount of lipid fractions was taken as 100%, the percentage of each lipid type being determined. In addition, we calculated the ratio of cholesterol fraction to total phospholipids (CS/PL). Microviscosity of macrophage membranes in lipid-lipid and protein-lipid interactions was estimated by recording lateral diffusion of a hydrophobic pyrene probe (Sigma-Aldrich, USA). An eximerization coefficient is inversely dependent on microviscosity and calculated by the formula

$$C_{ex} = \frac{F_{ex}}{F_{em}}$$

where $F$ is fluorescence at wavelength of 470 and 395 nm. Hydrophobicity degree was analyzed by the fluorescence of 1,6-diphenyl-1,3,5-hexatrien probe (Sigma-Aldrich, USA) [14]. Fluorescent spectra of membrane probes were recorded by a Fluorat-02-Panorama spectrofluorometer (Lumex, Russia).

During the second research stage we studied the functional state of peritoneal macrophages before and after exposure. Phagocytic activity was estimated by the absorption of 0.8-µm latex particles. We calculated the percentage of active cells (phagocytic index) and the average number of latex particles absorbed by a phagocyte (phagocytic number). Metabolic activity was assessed by the findings of spontaneous and zymosan-activated nitro blue tetrazolium tests (NBT test) [7]. The functional activity of peritoneal macrophages was determined by measuring luminal-dependent chemiluminescence using a Luminoskan Ascent microplate luminometer (Thermo Fisher Scientific, Finland). We carried out two parallel tests: spontaneous and induced by a stimulator (phytohemagglutinin, 200 µg/ml), the measurements being made within 30 min. We took into consideration the result at the peak of chemiluminescence and when its maximum was attained.

The results were presented as M±m, where M is arithmetic mean, and m is standard error of the mean. Mann–Whitney test was applied to determine the significance of differences. Two samplings were considered to belong to different general populations if p<0.05.

Results and Discussion

Structural condition of peritoneal macrophage membranes. At the first research stage we assessed a lipid spectrum of peritoneal macrophage membranes (Table 1). Spark plasma exposure for 60, 300, 600, and 1,200 s revealed a successive increase of CS ester (CSEs) fractions by 23.7–34.0%. The decrease in phosphatidylcholine fraction by 40.0–46.8% was found when exposure time was 300, 600, and 1,200 s. In addition, if exposure time is 1,200 s, there is a significant decrease in CS amount, by 14.7%, as well as the decrease of quantitative amount of such PL groups as phosphatidylethanolamine and lysophosphatidylcholine — by 49.0 and 60.4%, respectively. CSEs is the main form of CS supply in a cell, and the increase of their proportion can be related to possible oxidative reactions after spark plasma radiation, since the increase of total CS (CS+CSEs) in cells is known to be a protective response of a cell providing cell structural stability and protection against oxidation products [15, 16]. Phosphatidylcholine and phosphatidylethanolamine play a key role in membrane bilayer stabilization. The reduction of their amount can be related to the oxidation of certain PL fractions and result in the alteration of biophysical properties of cytolemma. Due to the fact that lysophosphatidylcholine exhibits the properties of a macrophage chemoattractant and mitogen, its reduction can have an effect on functional capabilities of a cell. Generally, the lipid spectrum changes at exposure time being 1,200 s suggests the switching cell protective reactions [15]. Moreover, the CS/PL ratio increases by 1.4 times, when being exposed within 1,200 s. It indicates the redistribution of relationship of the main lipid constituents and can result in the functional activity change of cell membranes.

UV radiation of a quartz lamp influences the lipid composition of peritoneal macrophage membranes to a greater extent than plasma radiation (See Table 1). 1,200-second exposure causes a significant reduction of lysophosphatidylcholine, phosphatidylcholine, and phosphatidylethanolamine by 53.2, 72.9, and 52.4%, respectively. CSEs proportion increases by 31.1%, the proportion of lysophosphatidylcholine by 53.2, 72.9, and 52.4%, respectively. CSEs is the main form of CS supply in a cell, and the increase of their proportion can be related to possible oxidative reactions after spark plasma radiation, since the increase of total CS (CS+CSEs) in cells is known to be a protective response of a cell providing cell structural stability and protection against oxidation products [15, 16]. Phosphatidylcholine and phosphatidylethanolamine play a key role in membrane bilayer stabilization. The reduction of their amount can be related to the oxidation of certain PL fractions and result in the alteration of biophysical properties of cytolemma. Due to the fact that lysophosphatidylcholine exhibits the properties of a macrophage chemoattractant and mitogen, its reduction can have an effect on functional capabilities of a cell. Generally, the lipid spectrum changes at exposure time being 1,200 s suggests the switching cell protective reactions [15]. Moreover, the CS/PL ratio increases by 1.4 times, when being exposed within 1,200 s. It indicates the redistribution of relationship of the main lipid constituents and can result in the functional activity change of cell membranes.

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The effect of gas discharged plasma radiation and UV radiation of a quartz lamp on the alterations in lipid spectrum of peritoneal macrophages is of oxidative nature, there is the decrease of PL, while CSEs is increasing. However, CS/PL ratio after quartz lamp
Table 1

Lipid composition of rat peritoneal macrophages after gas discharge exposure (%)

| Exposure type and time | LPC | SPH | PC | PE | FFA | CS | TAG | CSEs | CS/PL |
|------------------------|-----|-----|----|----|-----|----|-----|------|-------|
| **Before exposure**    | 2.05±0.20 | 7.39±0.27 | 3.65±0.39 | 2.06±0.32 | 13.74±1.44 | 26.79±0.87 | 5.32±0.61 | 35.66±2.83 | 1.87±0.10 |
| **Pilimin series IR-10** | | | | | | | | | |
| 30 s                   | 1.96±0.13 | 8.37±0.65 | 4.09±0.50 | 1.63±0.20 | 13.33±0.79 | 25.55±1.26 | 4.83±0.35 | 43.16±2.01 | 1.49±0.17 |
| 60 s                   | 1.61±0.13 | 7.37±1.07 | 3.76±0.37 | 1.69±0.16 | 11.29±0.42 | 25.00±1.08 | 4.76±0.52 | 44.13±0.94* | 1.99±0.34 |
| 300 s                  | 1.43±0.19 | 6.44±0.46 | 3.94±0.24* | 1.27±0.20 | 11.22±0.84 | 23.46±1.46 | 5.51±0.58 | 49.01±1.74* | 2.25±0.20 |
| 600 s                  | 1.65±0.20 | 7.21±0.52 | 3.17±0.10* | 1.42±0.18 | 13.62±1.44 | 24.65±1.44 | 5.65±0.55 | 45.87±2.25* | 2.19±0.13 |
| 1,200 s                | 0.81±0.19* | 6.12±0.51 | 3.04±0.10* | 1.05±0.10* | 13.17±1.67 | 22.84±0.87* | 4.68±0.37 | 47.82±3.32* | 2.58±0.23* |
| **DBK-9**              | | | | | | | | | |
| 30 s                   | 1.85±0.18 | 8.21±1.35 | 5.08±1.01 | 2.39±0.56 | 15.12±1.51 | 23.70±0.69 | 5.37±0.61 | 33.71±3.40 | 1.60±0.25 |
| 60 s                   | 1.43±0.28 | 8.50±1.60 | 4.59±1.06 | 2.44±0.56 | 14.99±0.94 | 24.06±1.28 | 5.03±0.33 | 37.06±2.98 | 1.71±0.48 |
| 300 s                  | 1.53±0.14 | 8.75±1.63 | 3.52±0.59 | 1.90±0.56 | 13.06±1.00 | 25.26±1.97 | 5.22±0.17 | 37.36±5.24 | 1.89±0.40 |
| 600 s                  | 1.60±0.14 | 9.45±0.32 | 3.72±0.82 | 2.31±0.15 | 14.06±0.88 | 25.47±1.12 | 4.38±0.44 | 35.75±2.94 | 1.80±0.22 |
| 1,200 s                | 0.96±0.08* | 5.74±0.99 | 0.99±0.06* | 0.98±0.13* | 14.52±1.25 | 26.69±0.49 | 3.86±0.34 | 46.74±1.47* | 3.23±0.43* |

Note: LPC: lysophosphatidylcholine, SPH: sphingomyelin, PC: phosphatidylcholine, PE: phosphatidylethanolamine, FFA: free fatty acids, CS: cholesterol, TAG: triacylglycerides, CSEs: cholesterol esters, CS/PL: the ratio of cholesterol fraction to total phospholipids; * significant difference with a control group (before exposure).

Table 2

Indices of membrane microviscosity of peritoneal macrophages after gas discharge exposure (relative units)

| Time exposure | Pilimin series IR-10 | DBK-9 |
|---------------|----------------------|-------|
|               | Lipid bilayer        | Protein-lipid interactions | Lipid bilayer | Protein-lipid interactions |
| **Unexposed** | 0.42±0.024           | 0.419±0.008               | 0.42±0.024 | 0.419±0.008 |
| 30 s          | 0.375±0.025          | 0.390±0.006* (p=0.036)    | 0.417±0.032 | 0.397±0.021 |
| 60 s          | 0.366±0.017          | 0.355±0.027* (p=0.013)    | 0.412±0.023 | 0.391±0.015 |
| 300 s         | 0.347±0.021          | 0.350±0.016* (p=0.016)    | 0.361±0.014 | 0.359±0.016* (p=0.010) |
| 600 s         | 0.330±0.015* (p=0.031) | 0.357±0.019* (p=0.022)   | 0.317±0.024* | 0.342±0.013* (p=0.019) |
| 1,200 s       | 0.261±0.016* (p=0.008) | 0.357±0.021* (p=0.043)   | 0.254±0.022* (p=0.004) | 0.322±0.017* (p=0.009) |

* Significant difference of values with a control group.

Condition of Peritoneal Macrophages after Gas Discharge Exposures
It is commonly known that membrane microviscosity increases in the area of lipid-lipid and protein-lipid interactions can result in the conformational alteration of membrane enzymes, intramembrane protein mobility and activity, impaired selective permeability and operation of cell receptors [18].

Fatty acids of PL provide plastic and flow properties of a bilayer. Therefore, in addition, to estimate the structural organization of fatty acids of PL (PL tails) we analyzed fluorescence of 1,6-diphenyl-1,3,5-hexatrien probe, which enables to study the condition of a hydrophobic hydrocarbon layer of cell plasma membranes. There was found the hydrophobicity decrease of fatty acids of phospholipid membranes after spark plasma radiation by 56.3 and 68.5%, respectively, if the exposure time was 600 and 1,200 s (See Figure).

1,6-diphenyl-1,3,5-hexatrien probe is known to be located in a hydrophobic area parallel to fatty acid chains. Hydrophobicity reduction can be related to the regularity decrease of hydrocarbon tails of PL, the formation of clusters and dynamic membrane defects resulting in the enhanced nonspecific cell permeability [19, 20].

Cell membrane hydrophobicity tends to increase in all the exposure modes selected after UV radiation of a quartz lamp. The exposure lasting 600 s, hydrophobicity was found to grow by 36.7%. Hydrophobicity increase can indicate the thickening of a phospholipid bilayer and the membrane permeability reduction [19, 20].

Spark plasma radiation and UV radiation of a quartz lamp result in oppositely directed alterations of hydrophobicity of phospholipid fatty acids. Membrane hydrophobicity decreases under gas discharged plasma radiation and increases under UV lamp radiation that can be related, firstly, to different mechanisms of the factors used, and secondly, spark plasma radiation is known to be pulsed and 400 times lower against UV radiation of a quartz lamp [17].

Cytoplasmic membrane is known to play a key role in determining normal functioning of both single organelles and a cell in whole. The alterations in membrane structural components can be called regulatory, since they determine functional condition of cells. The observed rearrangements in a lipid spectrum of peritoneal macrophages, microviscosity increase and the changes of membrane hydrophobicity can lead to both the limitation, as well as the enhancement of cell capabilities [2, 15].

Table 3

| Exposure time | Pilimin series IR-10 | DBK-9 |
|---------------|----------------------|-------|
|               | PI (%)    | PN     | PI (%)    | PN     |
| Before exposure| 53.43±1.23 | 3.52±0.06 | 53.43±1.23 | 3.52±0.06 |
| 30 s          | 54.14±1.14 | 3.69±0.15 | 54.80±1.19 | 3.58±0.17 |
| 60 s          | 57.88±1.13 | 3.68±0.15 | 55.33±1.69 | 3.50±0.11 |
| (p=0.034)*    |          |        |          |        |
| 300 s         | 54.63±1.39 | 3.70±0.12 | 52.67±1.86 | 3.48±0.11 |
| 600 s         | 53.02±1.12 | 3.83±0.13 | 55.25±2.50 | 3.66±0.14 |
| (p=0.040)*    |          |        |          |        |
| 1,200 s       | 50.17±1.66 | 3.47±0.14 | 55.20±1.02 | 3.51±0.12 |

Note. PI: phagocytic index, PN: phagocytic number; * significant difference of values with a control group.

Functional condition of peritoneal macrophages. When exposed to spark...
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Plasma radiation for 60 s, peritoneal macrophages were found to have a significant growth of phagocytic index by 8.3% (Table 3). The exposure time being 600 s, the phagocytic number increases by 8.8%. Phagocytic index growth suggests an increase of the number of cells participating in phagocytosis, and the phagocytic number increase indicates high absorbing capacity [1].

The analysis of the indices of metabolic activity of macrophages shows the activation of oxygen-dependent cell metabolism when exposed to spark plasma radiation for 300 s (Table 4). Generation intensity by reactive oxygen species cells is determined mainly by NADPH-oxidase activity, which depends on an initiating factor concentration and stimulation time [21]. UV radiation of a quartz lamp had no effect on phagocytic activity and the state of oxygen-dependent metabolism of peritoneal macrophages.

When estimating phagocytic activity by NBT test it is possible to record only absorption capacity of cells and the ability to produce superoxide anion radical, therefore, during the next stage we studied luminal-dependent chemiluminescence of cells, which enables to assess the formation of reactive oxygen species by macrophages including superoxide anion radical, singlet oxygen, hypochloride and general functional state of phagocytes. The experiment showed a significant increase of spontaneous chemiluminescence after spark plasma radiation exposure within 60 and 300 s by 65.3 and 116.5%, respectively (Table 5). If exposure time is 600 and 1,200 s, spontaneous chemiluminescence tends to decrease. There is no evident peak, though when a maximum level is achieved it decreases by 70.7 and 74.6%. 62.9–74.7% increase of spontaneous chemiluminescence was found under UV radiation of a quartz lamp, the exposure time being 300, 600, and 1,200 s. The time when maximum level of chemiluminescence occurs increases by 29.6% if exposure time is 1,200 s. Luminal-dependent chemiluminescence shows the activation of macrophages, which is due to the interaction of cells with reactive oxygen species formed as a result of radiation. Photobiological effects are considered to be implemented through a cascade of oxidative reactions, and an oxidative stress of lower intensity performs a regulatory function [22, 23].

To evaluate cell aptitude to respond on simulation, we analyzed the level of induced chemiluminescence. The exposure modes of 30, 60, and 300 s of spark plasma radiation were found to contribute to chemiluminescence activation by 1.6–3.9 times. When peritoneal macrophages were exposed to UV radiation within 300 and 600 s, induced chemiluminescence was observed by 1.3 and 1.4 times, respectively. Spark plasma radiation was shown to reduce the time of maximal chemiluminescence occurrence when UV radiation of a quartz lamp increases the reaction time at both spontaneous and induced forms.

### Table 4

#### Oxidative metabolism of peritoneal macrophages after gas discharge exposure

| Exposure time | Pilimin series IR-10 | DBK9 |
|---------------|-----------------------|------|
|               | Sp%       | Ind%  | Alsp         | Alind | MAC | RI | Sp%   | Ind%  | Alsp | Alind | MAC | RI |
| Before exposure | 53.25±0.48 | 57.25±1.31 | 0.61±0.02 | 0.69±0.03 | 1.92±0.02 | 1.09±0.03 | 53.25±0.48 | 57.25±1.31 | 0.61±0.02 | 0.69±0.03 | 1.92±0.02 | 1.09±0.03 |
| 30 s          | 54.0±1.15  | 57.33±2.85  | 0.65±0.02 | 0.70±0.03 | 1.98±0.01 | 1.03±0.01 | 51.0±1.29  | 56.25±1.49  | 0.58±0.02 | 0.65±0.02 | 1.90±0.01 | 1.12±0.02 |
| 60 s          | 54.50±1.32 | 58.0±0.71  | 0.69±0.03 | 0.74±0.01 | 1.95±0.02 | 1.05±0.02 | 53.80±2.35 | 57.60±2.25 | 0.62±0.03 | 0.68±0.03 | 1.92±0.01 | 1.09±0.01 |
| 300 s         | 57.67±2.19* | 60.33±2.03* | 0.71±0.02* | 0.77±0.02* | 1.97±0.03 | 1.07±0.04 | 51.50±2.02* | 55.75±1.03* | 0.58±0.01 | 0.64±0.01 | 1.91±0.02 | 1.09±0.02 |
| 600 s         | 54.67±2.33 | 59.67±0.33 | 0.69±0.06 | 0.75±0.05 | 1.93±0.03 | 1.08±0.04 | 51.67±1.45 | 54.0±1.47 | 0.56±0.03 | 0.62±0.02 | 1.89±0.03 | 1.10±0.02 |
| 1,200 s       | 53.0±0.58  | 59.3±3.18  | 0.67±0.02 | 0.73±0.03 | 1.90±0.04 | 1.13±0.05 | 51.40±0.51 | 56.40±0.81 | 0.59±0.02 | 0.66±0.01 | 1.90±0.01 | 1.11±0.02 |

**Note.** Sp%: spontaneous variant of NBT test, Ind%: induced variant of NBT test, Alsp and Alind: activity indices in spontaneous and induced variants, MAC: metabolic activity coefficient, RI: reserve index; * significant difference of values with a control group.
Thus, the oxidation of PL proceeds less intensively under spark plasma radiation. Membrane microviscosity in the area of lipid-lipid and protein-lipid interactions enhances to a greater degree after UV radiation of a quartz lamp. Hydrophobicity of PL of fatty acids enhances to a greater degree after UV radiation of a quartz lamp. In the area of lipid-lipid and protein-lipid interactions under spark plasma radiation. Membrane microviscosity decreases under spark plasma radiation and increases under UV radiation of a quartz lamp. Plasma radiation in contrast to UV radiation of a quartz lamp causes the growth of the number of cells participating in phagocytosis, the enhancement of absorbing capacity and oxygen-dependent metabolism of peritoneal macrophages. The period of an “oxygen explosion” of macrophages reduces after plasma radiation, and increases when exposed to UV radiation of a quartz lamp.

Conclusion. Spark plasma radiation compared to UV lamp radiation contributes to the enhancement of phagocytic activity of peritoneal macrophages in less continuous modes, the effects under study being more pronounced. The study findings enable to reveal optimal continuous modes, the effects under study being more functional cell activity after gas discharge exposure.

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### Table 5

**Luminal-dependent chemiluminescence in rat peritoneal macrophages**

| Exposure time | Pilimin series IR-10 DBK-9 Spontaneous chemiluminescence | Induced chemiluminescence | Pilimin series IR-10 DBK-9 Spontaneous chemiluminescence | Induced chemiluminescence |
|--------------|----------------------------------------------------------|----------------------------|----------------------------------------------------------|----------------------------|
|              | mV/min min                                              | mV/min min                 | mV/min min                                               | mV/min min                 |
| Before exposure | 0.170±0.029                                | 17.75±0.85                  | 0.223±0.014                                             | 9.38±0.80                  |
| 30s           | 0.199±0.037 *(р=0.010)*                           | 18.0±0.41                   | 0.359±0.049 *(р=0.001)*                                  | 7.0±1.13                   |
| 60s           | 0.281±0.029 *(р=0.043)*                           | 17.33±1.23                  | 0.817±0.137 *(р=0.001)*                                  | 9.33±1.54                  |
| 300s          | 0.368±0.054 *(р=0.037)*                           | 17.20±0.86                  | 0.866±0.096 *(р=0.001)*                                  | 12.67±1.50                 |
| 600s          | 0.086±0.021 *(р=0.019)*                           | 5.20±0.80                   | 0.243±0.047 *(р=0.0054)*                                 | 0.297±0.011 *(р=0.011)*    |
| 1,200s        | 0.079±0.023 *(р=0.030)*                           | 4.50±0.87                   | 0.174±0.014 *(р=0.004)*                                  | 0.277±0.028 *(р=0.030)*    |

* Significant difference of values with a control group.