Deep, large thermal burns are not limited to local lesions of tissues, they cause significant disruption of all systems and organs of the organism, change in metabolic processes. It is revealed that the primary links in the pathogenesis of burn disease are destruction of the skin, impaired neuroendocrine regulation and significant hemodynamic disorders. The reorganization of structures and impaired lung function, in response to a pathological process in the body, is attracting increasing attention of scientists. The aim of the study was to establish a submicroscopic rearrangement of the alveoli after a thermal lesion for 1 day after the experimental thermal trauma. Grade III burns were applied under ketamine anesthesia with copper plates heated in boiled water to a temperature of 97-100°C. The size of the lesion area was 18-20 % of the epilated surface of the body of rats. An experimental study of the structural components of lung alveoli after burn injury was performed on laboratory white male rats weighing 160-180 g. Euthanasia of rats was performed after ketamine anesthesia by decapitation. In the experiment, the study of the submicroscopic state of the walls of the alveoli of the lungs after thermal trauma was done. It is established that in the stage of shock after the application of burn injury - 1 day, in the alveoli of the respiratory department of the lung, there are adaptive compensatory and initial destructive changes of all structural components of the alveoli. Damage to the ultrastructure of the aerohematical barrier is manifested by intracellular edema and edema of the organelles of the endothelial cells, respiratory and secretory epitheliocytes, and the amount of heterochromatin increases in their deformed nuclei. The basement membrane also has signs of edema, sometimes homogeneous, fuzzy. The decrease in the number of vesicles and micropinocytotic vesicles in endothelial and respiratory epitheliocytes leads to impaired endothelial and alveolar metabolism. Numerous actively phagocytic alveolar macrophages with a well-expressed lysosomal apparatus are found in the alveoli. Initial alternative alterations of the ultrastructure of the components of the air-barrier barrier lead to disruption of gas exchange in the respiratory department of the lungs.

Keywords: lung alveoli, aerohematic barrier, submicroscopic changes, thermal trauma.

**Introduction**

Burns are the most common type of injury and are accompanied by significant changes in the structure and function of organs and systems of the affected organism [1, 3, 4, 5, 9, 11, 16]. In the early term after the injury disturbance of the enzyme homeostasis is caused by the disintegration of tissues in the lesion area, the subsequent destruction of the internal organs [10, 12, 18]. The disorders are directly related to changes in the burned skin as a source of biologically active substances that enter the bloodstream. Other sources of biologically active substances are organs and tissues that are not directly exposed to thermal effects, but are in a state of ischemia and circulatory hypoxia [3, 8, 13]. In the area of the lesion, after a thermal injury, a significant inflammatory reaction occurs, which is accompanied by the formation of biologically active substances, tissue breakdown products, specific and nonspecific toxins, which is the trigger mechanism of burn intoxication [8, 13, 15].

In the affected organism occurs centralization of circulation, which is adaptive in nature, but at the same time leads to significant disorders of regional and peripheral circulation. Not compensated disorders of microcirculation,
which accrue in time, lead to disorders of the systemic circulation and impaired function of all organs and systems [3, 7]. Hypovolemia is one of the leading factors leading to impaired lung respiratory function [14, 17, 19]. In a small circle of circulation the peripheral vascular resistance increases congestion develops. The decrease in circulating blood volume occurs mainly due to plasma loss [2]. Due to the loss of skin barrier function, as well as increased vascular permeability due to the action of vasoactive substances (histamine, serotonin, bradykinin, etc.), there is a pronounced plasma loss.

Given that the components of the respiratory department of the lungs after thermal trauma are not fully understood, it is advisable to conduct an in-depth study of the submicroscopic state of the cells of the alveoli wall of the respiratory department of the lungs.

Therefore, the purpose of the work was to establish a submicroscopic rearrangement of alveoli of the lungs of animals after thermal lesion in the shock stage after the experimental burn injury.

Materials and methods

The experiments were performed on 15 adult white male rats. The animals were kept on the standard diet of the vivarium of Ivan Horbachevsky Ternopil National Medical University of the Ministry of Health of Ukraine. Animal care and all manipulations were carried out in accordance with the provisions of the "European Convention for the Protection of Vertebrate Animals Used for Experiments and for Other Scientific Purposes" (Strasbourg, 1986) and in accordance with the provisions of the "General Ethical Principles for Experiments on animals", approved by the First National Congress on Bioethics (Kyiv, 2001). Grade III burns were applied under ketamine anesthesia with copper plates heated in boiled water to a temperature of 97-100°C. The size of the lesion area was 18-20 % of the epilated surface of the body of rats. On a daily examination, we monitored their general condition, the extent of local changes in the area of the burn wound, body weight, and mortality. The object of the study was the lungs. To study submicroscopic changes, animals were decapitated under ketamine anesthesia for 1 day, which, according to modern concepts, corresponds to the stage of shock of burn disease [13].

For ultrastructural studies, lung pieces were collected, fixed in 2.5 % glutaraldehyde solution, postfixed with 1 % osmium tetroxide solution on phosphate buffer. Further processing was carried out according to conventional methods [5]. Ultra-thin sections made on an LKB-3 ultramicrotome were counterstained with uranyl acetate, lead citrate according to the Reynolds method, and studied in an electron microscope PEM-125K.

Results

Electron microscopic studies have shown that the stage of shock for hemocapillaries is characterized by reactive changes, which are manifested by the enlargement of the lumen, significant blood filling and stasis formation. In endothelial cells, the initial signs of the destructive changes that are manifested by the disruption of the nucleus and cytoplasmic organelles are established. The hypertrophied nuclei have a rounded shape, with some of the invasions of karyolemma and the indistinct outlines of its membranes. Perinuclear space in some areas is expanded. The karyoplasm contains mainly euchromatin, however, in the part of the nuclei, marginally located tubercles of heterochromatin are noted. Enlarged tubules of the endoplasmic reticulum are located in the perinucleus zone, and the Golgi complex is represented by thickened cisterns and large vesicles. Peripheral, cytoplasmic areas of the endothelial cells were characterized by enlightenment and focal edema and a decrease in the number of pinocytosis vesicle and caveolae (Fig. 1).

The nuclei of the respiratory alveolocytes have a rounded
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Shape, fuzzy contours of the cariole membrane and invaginations. In moderately electron-dense karyoplasm, euchromatin prevails. The cytoplasm of cells is swollen, enlightened, especially in the peripheral regions. The integrity and structural organization of most organelles is partially impaired, with little pinocytosis vesicles. The basement membrane of the aerohematic lung barrier is characterized by swelling, thickening, or some areas of thinning (Fig. 2).

In the stage of burn shock in the secretory alveolocytes rounded nuclei are observed, membranes of the karyolemma are fuzzy, form shallow indentations, in the karyoplasm, euchromatin prevails, nucleolus are found. At some loci, the perinuclear space is enlarged and few nuclear pores are detected. In the cytoplasm of the secretory epitheliocytes, degranulation is observed, which is accompanied by a decrease in the amount of osmiophilic material in the lamellar bodies, and young forms are also present. The tubules of the endoplasmic reticulum are unevenly expanded, with few ribosomes on their membranes. The Golgi complex of type II alveolocytes is represented by expanded cisterns and a small number of vesicles and vacuoles. The mitochondria are hypertrophied, with electron-light matrix and fuzzy crystals. The apical surface of the cells contains few small microvilli (Fig. 3).

In the lumen of the alveoli, there is an increase in the number of alveolar macrophages that are in a state of increased functional activity. A considerable number of cytoplasmic outgrowths are found on their surface. Macrophage nuclei have irregularly shaped and invasive karyolemma. In the karyoplasm prevail euchromatin, there is one or two nuclei. The cytoplasm contains small, osmiophilic primary lysosomes and small secondary lysosomes. Small mitochondria have different shapes and their matrix is electron-dense. Dicystosomes of the Golgi complex are located mainly in the paranuclear space of the cytoplasm and are formed by expanded cisterns, small vesicles and vacuoles. The tubules of the granular endoplasmic reticulum are non-stretchable and thickened, with a moderate number of ribosomes on their membranes (Fig. 4, Fig. 5).

Discussion

Conducted submicroscopic studies of the components of the alveoli of the lungs in the stage of burn shock in thermal trauma revealed that the pathological link is impaired microcirculation in the walls of the alveoli, and causes the development of local hypoxia and increased proteolysis, permeability of the wall of microvessels with access to the alveoli of plasma components having lipolytic and proteolytic properties, which is fully consistent with scientific works [7, 13]. The expressed disturbances of microcirculation in the lungs during burn shock are accompanied by blockage of the capillary bed, stasis, bypass of blood flow, as well as violation of the aerohematic barrier. Interstitial edema creates hypoxia conditions for alveoli cells and, first of all, for large alveolocytes that produce surfactant.
In the stage of burn shock in the secretive alveolocytes in the cytoplasm there is degranulation, which is accompanied by a decrease in the number of lamellar bodies and osmiophilic material in them. Submicroscopically, there is an increase in the number of alveolar macrophages in the alveolar lumen that are in a state of increased functional activity.

During the period of shock after skin burns, pulmonary edema is detected as a violation of vascular permeability. Significant in this belongs to the segmented leukocytes. Violation of the permeability of microvessels develops as a result of the complex action of a number of factors that are in the blood and released locally by cellular elements during their degradation. The second component of the system of phagocytic cells, together with leukocytes, which are subject to urgent mobilization under conditions of burn shock are alveolar macrophages. Their increase is an adaptation reaction, a manifestation of the tension of all protective systems in a situation critical to the body [14, 17].

In the swollen areas of the lung, in the alveoli there is perfusion, but there is no aeration, which contributes to the strengthening of arterial hypoxemia in the period of shock. Thus, pulmonary edema is one of the links in the pathogenesis of burn shock.

However, the lungs are the only organ where not only blood circulation provides the vital function of the organ, but the organ itself exists for the circulation of the main function - gas exchange. Therefore, the unity of the functioning of the circulatory and respiratory systems is a necessary, adaptive pattern [13].

Thus, the results obtained from our own studies correlate with the scientific data of other researchers [13, 14, 17] and allow to conclude that severe thermal trauma in the early term of the experiment leads to disturbances of the ultrastructure of the aerohemal lung barrier. At the stage of shock in the experimental burns, adaptive compensatory changes and the initial signs of destructive ones were established. Hemocapillaries within the alveoli wall have enlarged blood-filled lumps, stasis formation is detected. Swelling of respiratory epitheliocytes and endothelial cells, disturbances in their cytoplasm of organelles, reduction of pinocytosis leads to the deterioration of gas exchange processes in the lungs.

Further studies plan to determine the degree of morphological changes in the structural components of the lung alveoli in dynamics after experimental thermal trauma under the condition of using corrective drugs.

Conclusions
Severe thermal injury leads to disruption of the ultrastructure of the aerohemal lung barrier. In the stage of shock during the experimental burns, adaptive-compensatory processes and signs of destructive changes of the alveolar epithelium, macrophages and the wall of the hemocapillaries are established, which leads to disruption of gas exchange processes in the lungs.

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Субмикроскопические изменения альвеол респираторного отдела легких через сутки после экспериментальной термической травмы

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Глубокие, велики за площадью термические ожоги не ограничиваются только местными урежениями тканей, они вызывают значительные поражения всей системы и органов организма, изменяют обменные процессы. Установлено, что первичными ланками к патогенезу ожоговой травмы являются разрушение кожного покрова, нарушение нейроэндокринной регуляции и значительные нарушения всех систем и органов организма. В стадии шока после нанесения ожоговой травмы - 1 сутки, в альвеолах респираторного отдела легких втягиваются известь термических ядер, иногда гемоглобин, в них образуются интрацеллюлярные и интерцеллюлярные отложения. Установлено, что первичными ланками у альвеолы легких, аэрогематический барьер, субмикроскопические изменения, термическая травма.

Ключевые слова: альвеолы легких, аэрогематический барьер, субмикроскопические изменения, термическая травма.

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