Introduction

In modern conditions of intensive industrialization and more increasing growing worldwide use of heat sources in everyday life, there is an increase in the frequency of burn traumatic injuries [10, 12, 13, 14, 18, 20]. Deep widespread burns are characterized not only by damage to the integumentary tissues, but also cause burn disease, which is characterized by different, lasting and peculiar general morphological and functional changes of all organs and systems of the organism [16]. It is clear that burn injury of the skin causes significant changes, first of all, skin cells [15], but damage to the skin is considered to be the main pathogenic mechanism that causes the development of burn disease, the components and factors of which are: generalized catabolic response in the trauma center and in all internal organs, systemic inflammatory and apoptotic responses, endogenous intoxication and multiple organ dysfunction [3, 8, 9].

Particularly unfavorable is the course of burn disease in the context of its association with diabetes mellitus [6, 16]. However, structural changes enterocytes of the mucous
membrane duodenum the conditions of association burn disease with diabetes have so far remained unheeded by researchers. On this basis, the important role of duodenal dysfunction in the pathogenesis of destructive and regenerative processes in its mucous membrane at different times after burn injury under the conditions of the course of diabetes mellitus.

The purpose of the work was to study the structural changes of enterocytes in the mucous membrane of the duodenum in burn injury of the skin of rats under the conditions of experimental streptozotocin-induced diabetes mellitus.

Materials and methods
This research was conducted on laboratory white adult male rats weight 180-210 g. The control group consisted of 21 animals without somatic pathology, the first experimental group consisted of 21 rats with burn skin injury, the second experimental group consisted of 21 rats with skin burn and experimental streptozotocin-induced diabetes. All studies and control of the animals were conducted in accordance with the rules for the use of the animals in the experiments, adopted by the "European Convention for the Protection of Vertebrate Animals Used for Experimental and other Scientific Purposes" (Strasbourg, 1986), "General Ethica Principles of Animal Experiments", adopted by the First National Congress On Bioethics (Kyiv, 2001), "Ethical Principles and Guidelines for Experiments on Animals: 3rd Edition" (Switzerland 2005) and the Law of Ukraine "On the Protection of Animals from Cruel Treatment" (2006). The experimental diabetes model was reproduced by administering streptozotocin to the rats intraperitoneally at a dose of 50 mg/kg, pre-dissolving it in 0.1M citrate buffer solution (pH-4.5). The duration of the experiment was 1 month. The control of the development of hyperglycemia in the second experimental group was blood glucose level - 24.24±0.79 mmol/L. In the control group 8.03±0.4 mmol/L.

In our research, burn skin injury was caused in accordance with the widespread model among researchers Regas F.C. and Ehrlich H.P. [17], which has been slightly modified and optimized Gunas I.V. with co-authors [7]. In the experimental simulation of skin burns, two copper plates in the form of an ellipse were kept in water at 100 ° C for 10 minutes and, under the conditions of ether anesthesia, were simultaneously applied symmetrically to both exposed rats with an exposure of 10 seconds. Burn skin damage in rats was IIA-B grade - dermal surface burn (according to the old classification IIIA grade) with a total area of 21-23% of the body surface with the development of burn shock. For morphological research the department of the duodenum was selected, a fragment of which was processed by conventional methods of light and electron microscopy. Semi-thin and ultrathin sections from epoxy blocks were obtained on an LKB ultratome (Sweden). Ultra-thin sections after appropriate contrast were examined under a PEM-125K electron microscope. Semi-thin sections were stained with methylene and toluidine blue. Sections of paraffin blocks were stained with hematoxylin-eosin.

The main criteria for assessing damage enterocytes of the mucous membrane duodenum were the results of histological and ultrastructural data in dynamics after 7, 14, and 21 days after skin burns. Within the specified time rats were injected a single intraperitoneally large dose of sodium thiopental and removed from the experiment by decapitation.

Results
In the duodenum mucous membrane of the animals of the first experimental group 7 days after burn found dystrophic changes in most enterocytes while maintaining the shape of unevenly swollen and infiltrated by lymphocytes (as well as leukocytes) villi (Fig. 1). Columnar borderless enterocytes, goblet cells, and intestinal...

Fig. 1. Leukocyte and lymphocytic infiltration of the swollen villi of the duodenal mucosa of the first experimental group 7 days after burn. Hematoxylin-eosin. x60.

Fig. 2. Ultrastructural organization of enterocytes with a brush border in the mucous membrane of the duodenum of the rat under normal conditions (control group of animals). The arrows indicate the inter-digitization of the cytoplasmic processes of adjacent enterocytes. 1 - brush border. EM x10000.
endocrinocytes were simultaneously altered. The described changes in the epithelial lining were accompanied by the expansion of the interepithelial spaces. These extensions revealed interepithelial lymphocytes, which also had pronounced damage to the structure. These morphological sings of the epithelial monolayer are significantly different from those under normal conditions.

In animals of the control group (Fig. 2), the apical portion of each columnar enterocyte with a brush border contains monofilaments, by means of which is formed dense intercellular contacts that isolate the lateral surface of enterocytes from the contents of the intestinal lumen. The cytoplasm of adjacent enterocytes has long thin processes that form interdigitations of complex shape along the contacting lateral surfaces of cells. Due to the availability of dense contacts and interdigitation of cytoplasmic processes, the maximum convergence of plasmalemma and intercellular spaces in the epithelium. Under the conditions of preservation of the integrity of the epithelial monolayer, a characteristic single-type, saved mainly leaf-like form of the villi of the mucous membrane duodenum.

In the mucous membrane of the duodenum of the animals of the first experimental group 7 days after burn, concomitant changes in the microcirculatory bed caused perivasal edema, leukocyte infiltration, diapedesis of erythrocytes, rupture of individual micro vessels with the formation of hemorrhages. In comparative analysis of the structural organization of the mucosa of the duodenum in this group of animals observed manifestations of adaptive processes. Detected the availability of a more pronounced folding of the mucous membrane. The villi are well developed, thickened, their own lamina includes a considerable number of lymphocytes.

Widespread phenomenon in this time interval is a variety of ultrastructural changes of enterocytes against the background of swelling of their cytoplasm (an indicator of which is the enlightenment of the cytoplasmic matrix): from the vacuolation of the tubules of the endoplasmic reticulum and damage to the mitochondria (manifestation is their intense swelling, fragmentation of the crust and inner membrane), to the complete destruction of organelles, the appearance of defects of plasmalemma and nuclear membrane.

In areas of enterocytes cytoplasm with partially lost brush border there are rounded autophagosomes and autophagolysosomes of different size with heteromorphic electron-dense contents (Fig. 3).

Fig. 3. Autophagosomes (1) of various sizes with heteromorphic electron-dense contents in the cytoplasm of enterocyte with the remnants of the brush border in the mucous membrane of the duodenum of the rat of the first experimental group 7 days after burn, 2 - the remnants of the brush border. EM x20000.

Morphological evidence of the initial stage of autophagy is the grouping of damaged cell organelles in certain loci of the cytoplasmic matrix and their sequestration by concentric coverage of the characteristic autophagosome structure - phagophore (double insulating membrane). In the future, autophagosomes merge with lysosomes and form autophagosomes with different electron density and structure (which is an indicator of the stages and efficiency of digestion of the sequestered material).

Digestion of the content of autophagolysosomes is accompanied by destruction of the inner membrane of the phagophore (under these conditions, the products of digestion are likely to be absorbed by the cytoplasm of the enterocyte). If part of the material remains undigested, then the autophagolysosome is transformed into an autophagic vacuole, which is directed to the plasmalemma of the apical region of the enterocyte and releases its contents outside.

In the animals of the first experimental group, erosion, ulcers, numerous small, and sometimes quite extensive hemorrhages were observed much more frequently than in the previous study period 14 days after the burn. In the population of the columnar enterocytes with the brush border increased the content of dystrophic altered cells,
Structural changes of duodenal mucosa enterocytes of rats in burn skin injury under experimental...

which were located not only in the upper, but also the middle and lower part of the villi.

The sections of the epithelial monolayer of enterocytes with preserved brush border alternated with areas free of brush border. Enterocytes of the epithelial monolayer were adjacent to desquamated cells of varying degrees of conservation (Fig. 4). In the apical part of most enterocytes with partially lost brush border were located groups of rounded autophagosomes and autophagolysosomes of different size and content (Fig. 5).

In the cytoplasm of many enterocytes with the presence of structural site defects of the plasmalemma and nuclear membrane in conjunction with local damage of the cytoplasmic matrix (which, given their variability, can probably be repaired) revealed signs of increased functional activity of organelles (evidence of moderate expansion of the tubules of the granular endoplasmic reticulum, an increase in the number of ribosomes, the presence of polyribosome, the aggregation of intact mitochondria, the integrity of the nucleus). In other enterocytes, at this time, the cytoplasm is vacuolated in the form of various enlargement of the tubules of the endoplasmic reticulum with enlightenment of their contents, vacuolar transformation of mitochondria, which is completed by complete necrotic destruction and cell fragmentation with the formation of cellular detritus.

In animals of the first experimental group after 21 days, after burns, morphologically focal atrophy of the mucous membrane was manifested by thickening and shortening of the villi; in some places they were completely absent. The erosions that were the result of enterocyte necrosis were often found (Fig. 6).

In animals of the second experimental group, subtotal necrosis of enterocytes with a brush border is widespread 7 days after burn, when, while maintaining part of the cytoplasm with an intact border, the cell area is subjected to necrotic degeneration, and the formed cell detritus with microvilli of the border enters the intestinal space.

14 days after burn in the mucous membrane of the duodenum of animals of the second experimental group, enterocytes in certain areas of the epithelial monolayer are subject to complete necrotic decay. In these areas, the basement membrane becomes "exposed" (Fig. 7). It is quite well preserved and even, it is somewhat thick compared to that of the norm (in the control group of rats).

The structural changes described above are accompanied by deformation and destruction of the villi of the duodenal mucosa (Fig. 8). The villi lose their typical leaf-like shape, often take on a "twisted" shape, and filamentous villi appear, surrounded by groups of desquamated enterocytes and cellular detritus.

21 days after burns in animals of the second experimental group in some parts of the mucous membrane of the duodenum of the crypt and villi are clearly deformed. The villi lose their typical cytoarchitectonics and have the appearance of clustered (and sometimes
today by researchers treated quite contradictorily [11]. According to modern data [1], autophagy is the process of digestion of its own distorted organelles and sections of the cytoplasm by lysosomes (thus autophagy is both a type of programmed cell death and a strategy for cell survival by recycling cellular material). Thus, the adaptive response of the duodenal mucosa enterocytes to the destructive effects of burn disease factors involves the use as part of the plastic and energy resources of the degraded cellular material to repair the damaged enterocyte and to maintain its viability. Under the conditions of our experiment in rats of the first experimental group, the course of structural changes of enterocytes of the duodenal mucosa is slow enough to include the adaptive mechanism of autophagy of distorted organelles. In the animals of the second experimental group, destruction of cytoplasmic and organelle enterocyte sites (due to the summation of the factors of burn disease and diabetes) is accelerated and cells die by necrosis.

Summarizing we can say that in the duodenal mucosa of the second experimental group, the dynamics of morphological changes during different periods after burns (7 days - the stage of shock and early toxemia; 14 days - the stage of late toxemia; 21 days - the stage of septicemia) differ from animals of the first experimental group. Comparison of the obtained data with the previously revealed ones suggests that the time intervals and the nature of adaptive changes of the duodenal mucosa are largely prolonged and worsened. Significant lesions of the mucous membrane should lead to disruption of the processes of the digestive system, parietal digestion and absorption, as well as immunological protection (considering that the mucous membranes are the first zone of contact of the body with the antigens of the environment and the leading link of immune protection, immune defenses), affects the condition of the body of burned persons and, to a large extent, determines the development of burn disease, as well as the course of diabetes.

The prospect of further research in this area is related to the study of the effects on the duodenum of drugs that reduce the intoxication of the body and blood sugar level.

**Conclusions**

The results of the studies showed that the base of damage to the duodenal mucosa enterocytes consist of deep destructive changes, which after 21 days (in the stage of septicotoxemia), as a rule, are not reverse and develop on the background of significant intoxication of the body. The structural changes of the enterocytes detected under the experiment were evidence of a violation of the structural integrity of the intestinal epithelial barrier. In animals of the first and (more) of the second experimental groups, the damaged epithelium of the duodenal mucosa critically weakens the adequacy of its function as an interface between the mucous membrane and the environment of the intestinal lumen, and is unable to structurally provide the reliability of antigenic toxicogenic bar.
In the mucous membrane of the duodenum in burn injury of the skin associated with diabetes is worsening of the manifestations of the adaptive reaction and prolongation of destructive processes, which is accompanied by a violation of the homeostasis of cellular interactions in the cytoarchitectonically altered and deformed crypts and villi.

References
дослідження показали, що в основі пошкоджень ентероцитів слизової оболонки двенадцятипалої кишки лежать глибокі деструктивні зміни, які через 21 добу (в стадії септикотоксемії), як правило, мають незворотній характер і розвиваються на фоні значної інтоксикації організму. При ожоговій травці шкіри, асоційованій з цукровим диабетом, в слизовій оболонці двенадцятипалої кишки погіршується прояв адаптивної реакції і пролонгуються деструктивні процеси, що супроводжується порушенням міжклітинних взаємодій в цитоархітектонічно змінених та деформованих ворсинках і криптах.

Ключові слова: ожогова травма, стрептозотоцин-індукований цукровий диабет, морфологічні зміни двенадцятипалої кишки.

СТРУКТУРНІ ЗМІНЕННЯ ЕНТЕРОЦИТІВ СЛІЗИСТОЇ ОБОЛОЧКИ ДВЕНАДЦАТИПЕРСТНОЇ КИШКИ ПРИ ОЖОГОВІ ТРАВМЕ КОЖИ В УСЛОВИЯХ ЭКСПЕРИМЕНТАЛЬНОГО СТРЕПТОЗОТИН-ИНДУЦИРОВАННОГО САХАРНОГО ДИАБЕТА

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Работа посвящена изучению структурных изменений энтероцитов в слизистой оболочке двенадцатиперстной кишки при ожоговой травме кожи крысы в условиях экспериментального стрептозотоцин-индуцированного сахарного диабета.

Исследования проведено на лабораторных белых половозрелых крысах-самцах массой 180-210 г. Группу контроля составили 21 животное без соматической патологии, первую экспериментальную группу составила 21 крыса с ожоговой травмой кожи, вторую экспериментальную группу составила 21 крыса с ожогом кожи у экспериментальным стрептозотоцин-индуцированным диабетом. Модель экспериментального сахарного диабета воспроизводили путем введения крысам стрептозотоцина внутрибрюшинно однократно в дозе 50 мг/кг. При экспериментальном моделировании ожога кожи две медные пластиники в виде эллипса выдерживали в воде при 100°С 10 минут и, в условиях эфирного наркоза, накладывали одновременно симметрично на обе обнаженные части тела крысы с экспозицией 10 секунд. Ожоговое повреждение кожи у крыс составляло IIА-Б степени - дермального поверхностного ожога (по старой классификации IIIА степень) общей площадью 21-23% поверхности тела с развитием ожогового шока. Для морфологических исследований изъяли отдел двенадцатиперстной кишки, фрагменты которого были исследованы общепринятыми методами световой и электронной микроскопии. Основными критериями оценки повреждения энтероцитов слизистой оболочки двенадцатиперстной кишки стали результаты исследования гистологических и ультраструктурных данных в динамике через 7, 14 и 21 сутки после ожога кожи. Результаты проведенных исследований показали, что в основе повреждений энтероцитов слизистой оболочки двенадцатиперстной кишки лежат глубокие деструктивные изменения, которые через 21 день (в стадии септикотоксемии), как правило, имеют незворотний характер и развиваются на фоне значительной интоксикации организма. При ожоговой травме кожи, ассоциированной с сахарным диабетом, в слизистой оболочке двенадцатиперстной кишки ухудшаются проявления адаптивной реакции и пролонгируются деструктивные процессы, сопровождающиеся нарушением межклеточных взаимодействий в цитоархитектонически измененных и деформированных ворсинках и криптах.

Ключевые слова: ожоговая травма, стрептозотоцин-индуцированный сахарный диабет, морфологические изменения двенадцатиперстной кишки.