ENDOGENOUS INTOXICATION IN ANIMALS OF DIFFERENT AGE GROUPS IN CASE OF POLYTRAUMA

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Background. Associated injury is a worldwide social and economic problem. Age related aspects of endogenous intoxication are not studied comprehensively. Annually, from 44 000 to 65 000 citizens die because of traumatic injuries. As a result, this number increased by 32.6% for the last 10 years. The detoxification system, as a component of the functional systems of the organism, experiences significant changes in case of polytrauma.

Objective. The study was aimed to discover pathogenetic peculiarities of the multiple trauma in age aspect in different disease periods and to explore the level of endogenous intoxication in this condition.

Methods. The experiments were performed on 72 white male rats aged 3, 6 and 12 months, which underwent simulation of severe skeletal trauma and examination of the contents of middle mass molecules and endogenous intoxication index (markers of endogenous intoxication) in 1, 4 and 24 hours after the associated injury.

Results. The most significant increase of the middle mass molecules was fixed in 24 hours after modeling of severe skeletal injuries in all groups of animals, especially it was the most pronounced in 12-month-old animals. The erythrocyte intoxication index reached the highest level in 4 hours after the injury, its increase was most significant in sexually mature adult animals.

Conclusions. A significant increasing of endogenous intoxication markers in 12-month-old rats, if compared to 3- and 6-month-old animals, can be caused by the decrease in compensatory protection mechanisms.

KEY WORDS: multiple trauma, endogenous intoxication, age.
were kept under the standard vivarium conditions. Decapitation was accomplished under thiopental anaesthesia on 1st, 4th, and 24th hour of the experiment in accordance to The European Convention for the Protection of Vertebrate Animals Used for Experimental and other Scientific Purposes (Strasbourg, 1986). We used blood serum for the research. The degree of toxic syndrome was evaluated by the level of middle mass molecules (MMM) [8] and erythrocyte intoxication index (EII) [10]. The received data was processed by method of statistical variation and Student t-test. The results were considered as significant at the value p<0.05.

**Results and Discussion**

In our experiment we detected the significant changes of MMM in all 3 animal groups and compared the results with the intact ones.

In the 3-month-old rats the increase of MMM was observed during the 1st hour after multiple trauma. The level of these molecules increased by 39%, and after 4 hours it was 48% if compared to the intact animals. On the 1st day after the multiple trauma MMM was on 43% higher than normal rates.

The similar changes were observed in the 6-month-old rats. During the 1st hour the MMM was increasing even by 73%, in 4 hours — by 83% and in 1 day of the experiment it was by 79% higher than in the intact group.

The most significant increase in MMM was evidenced in the 12-month-old animals. After the 1st hour it increased by 93%, in 4 hours — by 96% if compared to the intact animals, in 1 day — more than in 2 times.

The increase of MMM level in animals after multiple trauma proved the enforcement of catabolic processes. The rise of MMM levels, which can include oligopeptides, fragments of nucleic acids, fatty acids, and triglycerides, can prove the injury of the hepatocytes membranes and MMM that include purine bases, uric acid and aromatic amino acids — the suppression of the detoxifying function of liver [13].

The differences in the dynamics of changes of MMM between the 1st, 2nd and 3rd groups were manifested (Fig. 2).

After the multiple trauma the level of MMM in the 3-month-old animal group was higher by 82% if compared to intact animals. In 4 hours it was exceeding the level of control group by 108%, in 1 day — by 98%.
In the 6-month-old group MMM$_{280}$ was higher by 104% if compared to the control group, in 4 and 24 hours — by 136% and 98% respectively.

The highest level of MMM$_{280}$ was observed in the 12-month-old animals in all periods of the experiment. In 1 hour after multiple trauma, it was exceeding the level of the intact animals by 270%, in 4 hours — by 344%, in 1 day — by 448%.

We argue that the increase in MMM levels is a manifestation of catabolic processes enforcement in cells of vital tissues. The possible cause for these changes can be the activation of intracellular, particularly lysosomal proteases under the influence of these toxins [14].

The multiple trauma also influenced on the level of erythrocyte membranes impairment (Fig. 3).

EII significantly increased during the 1st hour after multiple trauma in the 3-month-old rats and exceeded the level of the intact animals by 85%. In 4 and 24 hours of the experiment it increased by 97% and 92% respectively.

In the 6-month-old rats, in 1 hour the EII was higher than in the intact animals by 107%, in 4 hours — by 124%, in 1 day — by 213%.

In the 12-month-old rats, in 1 hour the EII was higher by 127% if compared to the intact rats, in 4 hours — by 245%, and in 1 day — by 149%. In case of MMM the EII reached the highest level.

The data obtained prove the action potentiation of endogenous toxins that are excreted to blood in case of multiple trauma; it can be accompanied by either high level of catabolic processes or the suppression of the detoxification system functional activity [15].

**Conclusions**

MMM and EII proved serious age-related differences between the animals with multiple trauma. In all animal groups the highest value was reached during the 1st hour after the injury and continued with concomitant slow decline, especially in the 3- and 6-month-old animals in 4 hours after the injury. However, in the 12-month-old rats the increase of MMM and EII continued; it proved prolongation of endogenous intoxication.

The further research on endogenous intoxication will broaden the view on the pathogenesis of traumatic diseases in people of different age and in various periods of the diseases that will provide the opportunity to make prognoses on further course of the disease.

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