Bile is an important biological fluid, secreted by the liver and flows through the common bile duct into the duodenum. Bile contains bile acids (BAs), phospholipids, cholesterol, bilirubin, proteins, water and salts. The primary BAs are synthesized from cholesterol in the liver, conjugated to glycine or taurine to increase their solubility, secreted into bile, concentrated in the gallbladder during fasting, and expelled in the intestine in response to dietary fat, as well as bio-transformed in the colon to the secondary BAs by the gut microbiota, reabsorbed in the ileum and colon back to the liver. BAs in the intestine not only regulate the digestion and absorption of cholesterol, triglycerides, and fat-soluble vitamins, but also play a key role as signaling molecules in modulating epithelial cell proliferation, gene expression, and lipid and glucose metabolism by activating farnesoid X and G-protein-coupled bile acid receptor-1 in the liver, intestine, muscle and brown adipose tissue [1]. BAs receptors expressed in endothelial cells and may have important effects on both systemic and portal circulation [2]. The roles of BA in the pathogenesis of diabetes, obesity, metabolic syndrome, and cardiovascular diseases are seriously being considered, and BA and their derivatives seem to represent novel potential therapeutics to treat these diseases of civilization [3].

Obstructive (mechanical) jaundice induced by bile stones or malignancies preventing the normal outflow of bile and induces inexorably progressing hyperbilirubinemia with its consequent deleterious effects [4]. It requires a surgical treatment followed by external biliary drainage. The early and adequate decompression of bile ducts decreases postoperative complications, but lead to the external loss of bile [5]. The long-term loss of bile induces the disturbances in the digestion of lipids, in blood coagulation and other symptoms of vitamin A, B, D, E, and K deficiencies, in calcium metabolism disorder followed by bone osteomalation, disturbances in acid-base balance of blood and functions of the liver, kidneys and nervous system [6]. In our previous studies we found that 3-5 days of external biliary drainage (with full loss of bile) in rats induces the severe structural and metabolic disturbances in brain histaminergic neurons [7] as well as in neurons of parietal brain cortex [8]. The aim of the present paper was the estimation of histological changes in the cerebellum, in particular in its Purkinje cells, in rats in the similar setting of bile loss.

Material and methods

Animals, experimental design and chemicals

Experiments were performed on 40 male Wistar rats weighing 200±25 g. Rats were housed in vivarium with free access to standard laboratory food and kept under controlled environmental conditions. All experimental procedures complied with European Community Council Directive (86/609/EEC) for care and use of laboratory animals. This study was approved by the Biomedical Ethics Committee of the Grodno State Medical University.

In 20 rats the proximal part of common bile duct (3–5 mm below the confluence of the lobular hepatic ducts) was cut, ligated and all the bile secreted by the liver was collected through a polyethylene catheter into the outside glass bile container. The container was fixed to the skin on the right side of the rat body. 20 animals of the control group underwent sham operation (no removal of bile). On the 1th, 3th, and 5th days after the operation the animals of the control and experimental groups were decapitated and cerebellum cortex samples were collected for histology and cytochemistry.

Conclusion: Total loss of bile (on the 1st, 3rd, and 5th days) induces the gradual increase in structural and metabolic disturbances in cerebellar Purkinje cells and death of some of them.
for further cytochemistry. All the chemicals were obtained from Sigma-Aldrich (USA).

**Histology**

7 μm paraffin sagittal sections of the cerebellum cortex were stained with hematoxylin and eosin, 0.1% solution of thionine (the Nissl method) to assess general cytology of neurons and for the identification of dying neurons [9]. The examination of histological preparations, their microphotography and morphometry was carried out using microscope Axioskop 2 plus) equipped with digital camera Axiocam MRC5 (Carl Zeiss, Germany). In the preparations stained by the Nissl method the total amount of cerebellum Purkinje cells in the 1 μm interval of cortex gyrus was estimated, as well as the amount of normal and pathological types of neurons according to the intensity of their cytoplasm chromatophilia and the shape of cells bodies: normochromic (normal, medium staining), hyperchromic (intense staining), hyperchromic shrunken, hypochromic (pale staining) and shadow cells (very pale remnants of dead neurons).

Morphometry and cytophotometry of Purkinje cells in paraffin sagittal sections stained by the Nissl method were carried out using computer image analysis software Image Warp (Bit Flow, USA). To estimate the size and shape of neuronal bodies the images of up to 30 neurons bodies on the computer monitor were outlined by mouse cursor. Maximal and minimal diameter (D), perimeter (P), square (S), as well as form-factor (4πS/P2 – parameter of sphericity and folding) and the factor of elongation (maximal D/minimal D – parameter of sphericity) were calculated.

**Cytochemistry**

10 μm frozen sagittal sections of the cerebellum were prepared using cryostat (Leica CM 1840, Germany). The activity of the oxidizing enzymes, such as succinate dehydrogenase (SDH, EC 1.3.99.1), lactate dehydrogenase (LDH, EC 1.1.1.27), glucose-6-phosphate dehydrogenase (G-6-PDH, EC 1.1.1.49), NADH dehydrogenase (NADHDH, EC, 1.1.1.49), as well as the activity of marker enzyme of lysosomes acid phosphatase (AP, EC 1.4.3.4) were examined [10]. For the enzyme histochemistry the cryostat sections were placed into the corresponding incubation medium, including the buffer, substrate, co-factor, if necessary, and chromogen, for 30 min – 5 hours to visualize the location of enzymatic activity, then washed and embedded in the suitable plastic medium. The content of RNA in Purkinje cells cytoplasm was examined in paraffin sections by the Einarsson method [10]. The enzyme activities or products of histochemical reactions were determined in the cytoplasm of neurons on the optic density of chromogen obtained in the course of histochemical reactions. In each experimental group 150-200 neurons were estimated.

**Statistics**

The mean values obtained for every animal were processed with nonparametric statistics (because of the small number of animals in the groups) using software STATISTICA 10 (StatSoft, Inc., USA). In descriptive statistics, the values of median (Me) and interquartile range (IQR) were determined. The differences were considered significant at p<0.05 (Mann-Whitney U-test) because it was not a normal distribution.

**Results**

**Histology**

At the light microscopic level following 1 day of bile loss no structural changes in cerebellum Purkinje cells were found. Following 3 days of bile loss the shape of Purkinje cells became various: elongated, oval or triangle, but not pear-shaped only. The hyperchromic shrinking and dead neurons appeared. Following 5 days of bile loss the destructive changes in Purkinje cells increased. Highly elongated cells with extended apical dendrites and lyses of chromatophylic substance, or swelling of nucleus and cytoplasm, deformation and division of nuclei appeared (Fig. 1, 2). The single shadow cells were also observed. But there remained the cerebellum cortex regions where neurons were still unchanged.

**Figure 1. – The cerebellum Purkinje cells.** A – control (5 days after the sham operation); B, C – 5 days of bile loss (B – swelling of nucleus, C – division of nucleus). Nissl staining. Digital microphotography. Magnification x1000. Scale bars – 5 μm
The calculation of Purkinje cells have revealed that the amount of hyperchromic neurons on the 5th day of bile loss increased 3.8-fold (p=0.009), hyperchromic shrunken neurons increased 34.4-fold (p=0.009), a shadow cells – 7.5-fold (p=0.037). The amount of dead neurons estimated by the Victorov method increased 28-fold (p=0.018). In dynamics of bile loss the gradual decrease in Purkinje cells size, their elongation and loss of sphericity took place (Fig. 3).
**Histochemistry**

The loss of bile induces the gradual decrease in Purkinje cells cytoplasm, the activity of the mitochondria marker enzyme succinate- and NADH-dehydrogenase, as well as glucose-6-phosphate- and NADH dehydrogenase (NADHDH, EC, 1.1.1.49) and the activation of lactate dehydrogenase and the marker enzyme of lysosomes acid phosphatase (Fig. 4-7).

**Figure 4.** – Activity of succinate dehydrogenase in cerebellum Purkinje cells.
A – controls (5 days after sham operation); B – 5 days of bile loss (lower staining in cytoplasm). According to Nachlas et al. Digital microphotography. Magnification ×400. Scale bars – 10 µm

**Figure 5.** – Activity of lactate dehydrogenase in cerebellum Purkinje cells.
A – controls (5 days after sham operation); B – 5 days of bile loss (higher staining in cytoplasm). According to Hess, Scarpelli and Pearse. Digital microphotography. Magnification ×400. Scale bars – 10 µm

**Figure 6.** – Activity of acid phosphatase in cerebellum Purkinje cells.
A – controls (5 days after sham operation); B – 5 days of bile loss (higher staining in cytoplasm). According to Gomori. Digital microphotography. Magnification ×200. Scale bars – 10 µm
**Discussion**

In our previous paper following bile loss in rats we found similar dramatic structural and metabolic disturbances in brain histaminergic and parietal cortex neurons [7, 8]. The removal of bile from the body in the present study induces the increase in the number of hyperchromic shrunken Purkinje cells and shadow cells, which seems to be too severe for neurons to survive [11]. The decrease in size, loss of sphericity and elongation of neurons took place probably as a result of rise of osmolarity of intercellular fluid. The shrinkage of neurons occurs through the activation of Na⁺-K⁺-2Cl⁻ transport systems or coupled Na⁺/H⁺ и Cl⁻/HCO₃⁻ pathways of metabolism [11, 12]. The shape of neurons can be disturbed also by the damage of cytoskeleton.

One of the possible reasons of cell membranes damage and structural disturbances of neurons can be the loss of cholesterol, which is included into the composition of biological membranes and supplied by other organs to the liver for the synthesis of biologically important substances – bile acids, which are lost during the removal of bile. In addition, in the loss of bile and absence of its entering the duodenum, a long-term afferent impulsion from duodenum results in the process of central inhibition.

**Conclusion**

The loss of bile induces the gradual increase in structural and metabolic abnormalities in Purkinje cells resulting in severe, irreversible disturbances (hyperchromic shrunken neurons) and the death of neurons.
of some of them. A gradual decrease in Purkinje cells size, the loss of sphericity and elongation of neurons, followed by the activation of succinate-, NADH-, glucose-6-phosphate-dehydrogenases and activation of lactate dehydrogenase and acid phosphatase take place. All these changes reflect the ratio of the processes of damage and adaptation in the settings of bile absence in the body.

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References
КЛЕТКИ ПУРКИНЬЕ КОРЫ МОЗЖЕЧКА ПРИ ОТВЕДЕНИИ ЖЕЛЧИ У КРЫС

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Цель исследования – оценка структурных и метаболических изменений в клетках Пуркинье коры мозжечка крыс при полном наружном отведении желчи.

Материал и методы. Эксперименты выполнены на крысах-самцах Wistar, массой 200-250 г. У 26 крыс в области ворот печени в общий желчный проток вводили и фиксировали катетер из пластика, всю желчь, секретируемую печенью, собирали в стеклянную емкость, закрепленную снаружи. 20 животным контрольной группы провели ложную операцию (желчь не отводили). Через один, трое и пять суток после операции животных контрольной и опытной групп выводили из эксперимента, а образцы коры мозжечка забирали для гистологического и гистохимического исследования.

Результаты. Потеря желчи вызывала постепенное увеличение структурных и метаболических нарушений в клетках Пуркинье, что приводило к тяжёлым, необратимым нарушениям и даже гибели некоторых из них. Установлено постепенное уменьшение размеров перикарионов клеток Пуркинье, потеря их сферичности и увеличение большого радиуса, а также ингибирование сукцинат-, NADH-, глюкозо-6-фосфата-дегидрогеназ и, напротив, активация лактатдегидрогеназы и кислой фосфатазы, маркерного фермента лизосом. Все эти изменения отражают соотношение процессов повреждения и адаптации в поврежденных нейронах в условиях отсутствия желчи в организме.

Выводы. Полная потеря желчи в течение одних, трёх и пяти суток вызывает постепенное увеличение структурных и метаболических нарушений в клетках Пуркинье коры мозжечка и гибель некоторых из них.

Ключевые слова: мозжечок, клетки Пуркинье, структура, гистохимия, потеря желчи.

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