ACTIVITY OF THE LIVER MITOCHONDRIAL ASPARTATE AMINOTRANSFERASE AND MALATE DEHYDROGENASE IN RATS WITH TOXIC HEPATITIS UNDER CONDITIONS OF ALIMENTARY PROTEIN DEFICIENCY

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Nutritional demands in proteins depend on the life stage and health status of organisms. Both humans and experimental animals under stress conditions and especially drug processing become to be more sensitive to the protein deficit in the food. In this study, we examined some acetaminophen-induced metabolic effects potentiated by alimentary protein deprivation (APD) in rat liver. In particular, activities of the liver mitochondrial aspartate aminotransferase and malate dehydrogenase in rat liver were studied in conditions of balanced and imbalanced by protein diets of isocaloric content. It has been found that acute acetaminophen-induced hepatitis in comparison to control does not change the activity of mitochondrial malate dehydrogenase causing simultaneous 4-fold reduction in activity of mitochondrial aspartate aminotransferase and 2.5-fold reduction of mitochondrial oxaloacetate content. Interestingly, alimentary protein deprivation enhances the effects of acetaminophen on the described parameters. Finally, in order to confirm these associations between amount of the protein in the rat diet and physiological measures in their liver with toxic injury, principal component analysis (PCA) was performed. Two principal components characterize changes in physiological measures in our study. Principal component 1 explains about 86 % of the variation among whole dataset mainly related to control group and group subjected to acetaminophen treatment with simultaneous APD. It reveals the tight association of scores for AST activity and oxaloacetate level with control group, which might indicate the high efficiency in the oxaloacetate conversion by AST lacked in both groups with hepatitis. Similarly, principal component 1 explaining the variance in MDH activity shows its linkage to the control group, indicating the importance of MDH for the health status of control animals. On the other side, principal component 2 reveals close association between lactate and pyruvate levels as well as cytosolic NAD+/NADH ratio with acetaminophen-treated group of animals subjected to APD, confirming that toxic liver injury associated with low protein consumption leads to increased lactate-pyruvate turnover in cytosol affecting. This potentially might be associated with energy-generating dysfunction in liver under toxic hepatitis on the background of dietary protein deficiency.

Keywords: ACETAMINOPHEN, LIVER, INJURY, ASPARTATE AMINOTRANSFERASE, MALATE DEHYDROGENASE

АКТИВНІСТЬ МІТОХОНДРІАЛЬНИХ ЕНЗИМІВ АСПАРТАТАМІНОТРАНСФЕРАЗІ ТА МАЛАТДЕГІДРОГЕНАЗИ У ПЕЧІНЦІ ЩУРІВ З ТОКСИЧНИМ ГЕПАТИТОМ ЗА УМОВ АЛІМЕНТАРНОЇ НЕСТАЧІ ПРОТЕЇНУ

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Потреба у протеїнах залежить від стадії життя і стану здоров’я організмів. Як люди, так і експериментальні тварини в умовах стресу та особливо використання ліків стають чутливішими до дефіциту протеїну в їжі. Метою роботи було дослідження деяких метаболічних ефектів у печінці за умов нестачі харчового протеїну у тварин з ацетамінофен-індукуваним гепатитом. Зокрема, визначали активність мітохондріальніх ензимів аспартатамінотрансферази та малатдегідрогенази у печінці щурів з токсичним гепатитом за умов різної забезпеченості рацию харчовим протеїном. Встановлено, що за умов зростання ацетамінофен-індукуваного гепатиту не спостерігаються зміни активності мітохондріальної малат-
дегідрогенази при одноразовому 4-кратному зниженні активності мітохондріальної аспартатамінотрансферази та 2,5-кратному зниженні активності мітохондріального оксалоацетату. Ця відома, що альтернативна депривація протеїну посилює вплив ацетамінофену на описані параметри. Для того, щоб підтвердити залежність між кількістю протеїну в раціоні тварин і метаболічними змінами у печінці за умов токсичного ушкодження, було застосовано метод головних компонент (PCA). Дві головні компоненти характеризують зміни у нашому дослідженні. Головна компонента 1 пояснює близько 86 % відхилень між контрольною групою і групою протеїн-дефіцитних тварин з ацетамінофен-індукуваним ушкодженням. Виявлено цілу асоціацію балів для активності АСТ та рівня оксалоацетату порівняно з контрольною групою в обох групах з гепатитом. Аналогічно, головна компонента 2, що пояснює дисерцію в активності малитедегідрогенази, показує на важливість малитедегідрогенази для стану здоров'я тварин. За іншого боку, головна компонента 2 виявляє тісний зв'язок між рівнями протеїну і пірвовату, а також з активністю цитозольного NAD+/NADH у групі протеїн-дефіцитних тварин, які отримували ацетамінофен. Встановлений факт підтверджує, що окислочовокисний баланс печінки на підставі змін у складі протеїну супроводжується інтенсифікацією перетворення лактат-пірвовату у цитозолі. Це потенційно може бути пов'язано з дисфункцією енергозабезпечення у печінці при токсичному гепатиті на підставі альтернативної нестачі протеїну.

Ключові слова: АЦЕТАМИНОФЕН, ПЕЧІНКА, УШКОДЖЕННЯ, АСПАРТАТАМИНОТРАНСФЕРАЗА, МАЛАТДЕГІДРОГЕНАЗА

АКТИВНОСТЬ МИТОХОНДРИАЛЬНЫХ ЭНЗИМОВ АСПАРТАТАМИНОТРАНСФЕРЫ И МАЛАТДЕГИДРОГЕНАЗЫ В ПЕЧЕНИ КРЫС С ТОКСИЧЕСКИМ ГЕПАТИТОМ В УСЛОВИЯХ АЛИМЕНТАРНОЙ БЕЛКОВОЙ НЕДОСТАТОЧНОСТИ

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Потребность в белках зависит от стадии жизни и состояния здоровья организмов. Как люди, так и экспериментальные животные в условиях стресса и особенно при употреблении лекарств становятся более чувствительными к дефициту белка в пище. Целью работы было исследование некоторых метаболических эффектов дефицита пищевого белка в печени у животных с ацетаминофен-индуцированным гепатитом. Изучали активность митохондриальных ферментов аспартатаминотрансферазы и малатдегидрогеназы в печени крыс с токсическим гепатитом в условиях различной обеспеченности пищевым белком. Установлено, что в условиях остrego ацетаминофен-индуцированного гепатита активность митохондриальной малатдегидрогеназы не изменяется при одновременном 4-кратном снижении активности митохондриальной аспартатаминотрансферазы и 2,5-кратном снижении содержания митохондриального оксалоацетата. Интересно, что депривация пищевого белка усиливает действие ацетаминофена на описанные параметры. Для подтверждения зависимости между количеством протеина в пищевом рационе крыс и метаболическими изменениями в печени в условиях токсического повреждения, проведен анализ главных компонент (PCA). Две главные компоненты характеризуют изменения в нашем исследовании. Главная компонента 1 объясняет около 86 % отклонений между набором данных, связанных с контрольной группой и группой протеин-дефицитных животных с ацетаминофен-индуцированным повреждением. Установлена ассоциация баллов активности АСТ и уровня оксалоацетата по сравнению с контрольной группой у обеих групп с гепатитом. Аналогично, главная компонента 1, характеризующая дисперсию в активности малатдегидрогеназы, свидетельствует о важности малатдегидрогеназы для состояния здоровья. С другой стороны, главная компонента 2 обнаруживает тесную связь между уровнями лактата и пирувата, а также соотношением цитозольного NAD+/NADH в группе протеин-дефицитных животных, получавших ацетаминофен. Установленный факт подтверждает, что токсическое повреждение печени на фоне низкого употребления протеина сопровождается интенсификацией перераспределения лактат-пирувата в цитозоле. Указанные изменения потенциально могут быть связаны с дисфункцией энергообеспечения в печени при токсическом гепатите на фоне алиментарного дефицита белка.

Ключевые слова: АЦЕТАМИНОФЕН, ПЕЧЕНЬ, ПОВРЕЖДЕНИЕ, АСПАРТАТАМИНОТРАНСФЕРАЗА, МАЛАТДЕГІДРОГЕНАЗА
Proteins are essential macronutrients used in various cellular processes. A lack of protein in the diet (alimentary protein deficiency, APD) can cause a set of metabolic abnormalities associated with repressed growth and increased mortality [17]. At the organismal level, an important role in the coordination of protein metabolism belongs to the liver. Liver is also known to be the main detoxification organ during xenobiotic and drug administration [6]. As the processes of xenobiotic metabolism require multiple biochemical transformations, and some intermediates mediate toxic responses [5], one may suggest that liver might be potentially susceptible to drug injury upon the lack of protein in the diet. Such a case might be a problem in developing countries, where low dietary protein associated with high levels of pollutants and xenobiotics in the environment as well as administration of untested or old generation drugs in the medicine is a common issue. For example, acetaminophen (known also as paracetamol), one of the most widely used analgesics in the therapeutic practice in developing countries, possesses a high risk to cause liver injury [8]. However, the effects of acetaminophen administration associated with low protein intake on metabolic processes liver cells remains unknown.

In this study, we have examined some acetaminophen-induced metabolic effects potentiated by alimentary protein deprivation (APD) in rat liver. In particular, the effects of acetaminophen on cytoplasmic NAD+/NADH ratio and functioning of malate-aspartate shuttle mitochondrial enzymes in rat liver were studied in conditions of balanced and imbalanced by protein diets of isocaloric content.

**Materials and methods**

**Experimental Design and Procedures.** In the study, 8–10 week old white nonlinear rats weighing 90–100 g were used. Animals were kept in individual plastic cages with sand bedding; they were fed twice per day having *ad libitum* access to water. The experiment was conducted in accordance with the rules set by the “European convention for the protection of vertebrate animals used for experimental and other scientific purposes” (Strasbourg, 1986).

For the study, animals were divided into three groups: I — control group of rats maintained on the balanced diet (C, control group); II — rats with acute acetaminophen-induced hepatitis, maintained on the balanced diet (H, group with hepatitis); III — rats with acute acetaminophen-induced hepatitis, maintained under the conditions of alimentary deprivation in protein (APD+H, alimentary deprivation in protein + hepatitis). Each experimental group had from 6 to 10 animals. Over 4 weeks, animals of the C and H experimental groups were fed balanced diet containing 14 % protein (in the form of casein), 10 % fat and 76 % carbohydrates [4]. Animals of the APD+H group were fed for 4 weeks isoenergetic food containing 4.7 % protein, 10 % fat and 85.3 % carbohydrates. After 4 weeks, toxic hepatitis was caused in the H and APD+H experimental groups by *per os* administration of acetaminophen. For this, acetaminophen was added to the food at a dose 1 g/kg in a 2 % starch suspension for 2 days [14]. Animals were euthanized on the 31st day of the experiment. The rats were sacrificed by decapitation 48 hours after the paracetamol application, their livers were quickly removed.

Liver tissue was homogenized in 9 volumes of cold buffer at 4 °C. The homogenates were then centrifuged at 4 °C (1000 g/min, 10 min).

The mitochondrial fraction of the liver homogenate was isolated by differential centrifugation (*Heraeus Biofuge*, Germany) at 0–3 °C in the following buffer medium: 250 mM sucrose, 1 mM EDTA, 10 mM Tris-HCl; pH 7.4 at 0–3 °C. The cytosol fraction of homogenates was isolated after the separation of microsomal fraction [11].

**Enzyme assays.** The activities of mitochondrial aspartate aminotransferase (AST, EC 2.6.1.1) and malate dehydrogenase (MDH, EC 1.1.1.37) were measured spectrophotometrically at 340 nm as described earlier [2, 19].

**Lactate and pyruvate assays.** The content of lactate was evaluated using the enzymatic method according to [3] in the presence of lactate dehydrogenase — EC 1.1.1.27 (the final activity in the incubation mix — 2 IU/ml), 0.05 M NAD in glycine-hydrazine buffer (0.4 M hydrazine sulphate, 1 M glycine, 0.2 % EDTA-Na, pH 9.5) The formation of reduced NADH, which quantity is equivalent to the amount of oxidized lactate, was registered photometrically at the wavelength 340 nm.

The content of pyruvate in the cytosolic fraction was determined spectrophotometrically in test with iron (III) nitrate [7].
Oxaloacetate concentration measurement

Concentration of oxaloacetate was determined by colorimetric method based on the interaction of oxaloacetate 2.4-dinitrophenylhydrazine and formation of oxaloacetate hydrazone with maximum absorption at 546 nm [9].

The cytoplasmic NAD\(^+\)/NADH ratio assay

The cytoplasmic ratio NAD\(^+\)/NADH was calculated taking into account the equilibrium constant of the lactate dehydrogenase reaction as was proposed previously [13, 16]:

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\frac{\text{NAD}^+}{\text{NADH} + H^+} = \frac{1}{1.11 \times 10^{-4}} \times \frac{[\text{Pyruvate}]}{[\text{Lactate}]} \]

Statistical analysis. Statistical analysis was performed by T-test using Statistica 6.0 software (StatSoft, USA). All of the data was tested for normal distribution using Levene’s test and one-sample Kolmogorov-Smirnov test, respectively. Characteristics of the study group were expressed as Mean±Standart Deviation for normal distribution. The probability, P≤0.05 was accepted to have critical level of significance. Principal component analysis was done using JMP Pro 11 software (SAS Institute, USA).

Results and discussion

The hepatocytes’ system of energy bio-transformation determines the ability of an organism to restore its wellness after a toxic injury of many xenobiotics including acetaminophen [1, 12]. Simultaneously, the efficiency of hepatocytes to rebuild the metabolism upon xenobiotic stress depends on nutritional conditions. In this study, we aimed to disclose the biochemical adaptation of hepatocytes’ malate-aspartate shuttle mitochondrial enzymes in rats subjected to low protein diet upon acetaminophen-induced toxicity. The choice of malate-aspartate shuttle as an indicator of hepatocyte functional state is related to its role in NAD oxidoreduction. In particular, the malate-aspartate shuttle system is the central metabolic pathway to transfer glycolytic NADH from the cytoplasm to mitochondria for further NADH oxidation [15, 18].

First, we evaluated the activities of mitochondrial NAD-dependent malate dehydrogenase (MDH) and aspartate aminotransferase (AST). We have found that acetaminophen does not affect activity of mitochondrial NAD-dependent MDH in the liver of rats fed a protein balanced diet (fig. 1A). However, conversion of malate to oxaloacetate by this enzyme to be lower in the liver of animals subjected to APD. Thus, activity of malate dehydrogenase was 52 and 34 % lower in the APD+H group. Reduction in activity of mitochondrial aspartate aminotransferase (fig. 1B)

Fig. 1. NAD\(^+\)-malate dehydrogenase (MDH, A) and aspartate aminotransferase (AST, B) activities of liver mitochondria under toxic hepatitis with and without alimentary protein deprivation. The data are presented as means ± standard error means (S.E.M.), n = 8–10

Note: C bar shows the value of parameter in the liver of control group of rats maintained on the balanced diet; H bar shows the value of parameter in the liver of rats with acute acetaminophen-induced hepatitis, maintained on the balanced diet; APD+H bar shows the value of parameter in the liver of rats with acute acetaminophen-induced hepatitis, maintained under alimentary protein deprivation (APD). * — significantly different from the control (C) group. ** — significantly different from the group with acetaminophen-induced hepatitis. P≤0.05 represents statistical significance.
was found upon acetaminophen treatment when compared to the enzyme activity in liver of rats in the control group. However, APD enhanced the reduction in activity of mitochondrial aspartate aminotransferase in rat liver. Thus, the activity of the enzyme was 6-fold lower in the liver of ADP animals with acetaminophen-induced hepatitis than in the control group (fig. 1B).

Neither lactate nor pyruvate concentration were affected by acetaminophen in the cytosolic fraction in liver of rats (Fig. 2.A, 2B). However, the concentration of lactate and pyruvate in the cytosolic fraction of liver of rats subjected to APD with simultaneous acetaminophen treatment was 60 % and 3.3-fold higher respectively than in controls and animals with toxic hepatitis. In the opposite, the amount of oxaloacetate in liver mitochondria of rats decreased upon acetaminophen treatment. Moreover, APD enhanced the effect of acetaminophen (fig. 2C).

Since oxaloacetate in mitochondria is a substrate for both aspartate aminotransferase and citrate synthase, reduction in the activity of mitochondrial aspartate aminotransferase may indicate enhanced use of oxaloacetate in the tricarboxylic acid cycle (TCA) in the liver of acetaminophen-treated animals. It seems that APD enhances the use of oxaloacetate in liver mitochondria upon acetaminophen-induced hepatitis, which might lead to the increased oxidation of NADH formed in TCA by mitochondrial respiratory chain. As the state of energy supply to the cells primarily depends on the efficiency of the mitochondrial respiratory chain [10, 14], the reduction in activity of malate-aspartate shuttle mitochondrial enzymes in rat liver may indicate different rate of NADH oxidation in the cytosol and mitochondria.

Finally, in order to confirm these associations between amount of the protein in the rat diet and physiological measures in their liver with and without toxic injury, principal component analysis (PCA) was performed. Scree plot for the used types of diets and score plot for the measured metabolic parameters were combined in biplot for graphical representation of PCA (fig. 3). Two principal components characterize changes in physiological measures in our study. Principal component 1 explains about 86 % of the variation among whole dataset mainly related to control group and group subjected to acetaminophen treatment with simultaneous APD.
It reveals the tight association of scores for AST activity and oxaloacetate level with control group, which might indicate the high efficiency in the oxaloacetate conversion by AST lacked in both groups with hepatitis. Similarly, principal component 1 explaining the variance in MDH activity shows its linkage to the control group, indicating the importance of MDH for the health status of control animals. On the other side, principal component 2 reveals close association between lactate and pyruvate levels as well as cytosolic NAD\(^+\)/NADH ratio with acetaminophen-treated group of animals subjected to APD, confirming that toxic liver injury associated with low protein consumption leads to increased lactate-pyruvate turnover in cytosol affecting, thus, NAD\(^+\)/NADH flux.

**Conclusion**

The data suggests that alimentary protein deficiency might aggravate the metabolic imbalance in the liver after its toxic injury. Particularly, the association of toxic hepatitis with APD in rats leads to the reduction in the activities of malate-aspartate shuttle mitochondrial enzymes and glycolytic NADH inflow to the mitochondria. This potentially might be associated with energy-generating dysfunction in liver under toxic hepatitis on the background of dietary protein deficiency.

**Perspectives of the future investigations.**

The obtained results open prospects for the further study of the mechanisms of energy supply disturbance in the liver under conditions of its toxic damage with the different content of protein. It plan to investigate the activity of the main enzymes of glycolysis and the Krebs cycle in the liver under the experimental conditions for the purpose of justification of approaches to correction of energy disturbance.

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