To date, the diagnosis and treatment of extrahepatic cholestasis (EHC) at various stages of development remains one of the most pressing problems of modern biliary surgery. The purpose of the study is to determine the structural changes of the liver in patients with EHC of non-tumoral genesis according to shear wave elastometry and ultrasound in comparison with biochemical indicators of endotoxemia, inflammation and markers of fibrosis. Preoperatively, 121 patients with EHC of non-tumoral genesis were examined: standard general clinical studies, ultrasound examination of the liver and its shear wave elastometry were performed. Depending on the presence of jaundice and symptoms of hepatocyte damage, all the patients were divided into 4 groups. Serum content of medium molecular weight peptides, free hydroxyproline and glycosaminoglycans was determined. Statistical processing of the obtained data was performed using SPSS 16.0. In patients with EHC of non-tumoral genesis according to shear wave elastography, the stiffness of the liver varied depending on the increase in the intensity of jaundice. It was found that the serum bilirubin level in such patients was significantly different depending on the available jaundice and symptoms of hepatocyte damage both in comparison with the control group and between the study groups. In EHCs, structural changes in the liver depend on abnormalities in the biochemical composition of serum, which are significantly altered by prolonged extrahepatic cholestasis and manifested by severe liver failure. Thus, an increase in endotoxemia (p<0.001) was found in patients with EHC of non-tumoral genesis without jaundice and evidence of cholestasis development; indicators of liver stiffness depend on the level of bilirubinemia (r = 0.84), and in patients with hepatocyte damage also on the duration of cholestasis (r = 0.68). Bilirubin levels cannot be an indicator of long-term EHC.

Keywords: extrahepatic cholestasis of non-tumor genesis, elastometry, markers of endogenous intoxication, damage to hepatocytes.
noted that over the past 10 years, there has been a steady trend of increasing incidence of cholelithiasis, which is accompanied by the development of extrahepatic cholestasis [2, 3, 4, 5, 10].

Functional EHC is most often caused by Oddi sphincter dysfunction. Obstructive EHC, or mechanical jaundice, is a persistent violation of the excretion of bile from the bile ducts and gallbladder into the lumen of the duodenum, which can be caused by obstruction of the bile ducts by gallstones, helminths or mechanical compression of the bile ducts from the outside by tumor (pancreatic head, large duodenal papilla, common bile duct, enlarged gallbladder), inflammatory swelling of the mucous membrane of the biliary passages (non-purulent, sclerosing or purulent cholangitis), bile duct stenosis and other factors. Violation of the outflow of bile leads to its stagnation, increasing the pressure in the bile capillaries, their overstretched, increasing the permeability of the walls and their rupture, which promotes the flow of bile directly into the blood. The appearance of bile in the blood leads to the development of cholemic syndrome, and with complete obstruction of the bile ducts and the cessation of bile flow to the intestine (clinically, this is manifested in the acholic stool).

At detection of a syndrome of a cholestasis at the patient, for doctor it is necessary to make qualitative differential diagnostics with precise determination of degree and place of formation of the block and it is unacceptable to establish the diagnosis without excluding the possible causes of cholestasis. Diagnostic errors that occur in 12-38% of cases of cholestasis [3] lead to loss of precious time, occurrence of liver failure or other serious complications (gastrointestinal bleeding, purulent cholangitis, liver abscesses, sepsis) that in 14-27% of observations lead to death [1, 3, 8, 9, 10].

To date, prevention and treatment of EHC at various stages of development remains one of the most pressing problems of biliary surgery. There is evidence [11, 12, 13] that EHC affect the elasticity of the liver, but to date, the possibilities of using elastography to assess the functional and morphological status of the liver in EHC of non-tumorigenesis have not been sufficiently studied. Recently, several ultrasound and radiological methods have been developed to quantify liver elasticity. One of the most accurate and common among them is the method of shear wave elastography (SWE).

The basis of the method of shear wave elastography is the property of the ultrasonic beam to generate transverse mechanical shear waves in the direction of its propagation. The speed of their passage through the fabric depends on its stiffness or visco-elastic properties. The digital expression of liver stiffness is determined in kilopascals (kPa) from a specific control volume of the liver parenchyma [11-13]. However, the effect of EHC on liver elasticity in comparison with non-invasive serum markers of endotoxemia, inflammation, and fibrosis has not been studied.

Purpose of the study: to determine structural changes of the liver according to shear wave elastometry and ultrasound in comparison with biochemical indicators of endotoxemia, inflammation, markers of fibrosis in patients with non-hepatic cholestasis of non-tumorigenesis.

Materials and methods
The data of the examination of 121 patients with EHC of non-tumoral genesis who were undergoing treatment at the department of digestive surgery of the State Institution "Institute of Gastroenterology of the National Academy of Medical Sciences of Ukraine" for the period from 2013 to 2019 were analyzed.

Extrahepatic cholestasis in 22 patients (18.18%) patients was caused by chronic calculous cholecystitis with the phenomena of Oddi sphincter, in 28 patients (23.14%) - choledocholithiasis after cholecystectomy, in 18 patients (14.87%) - stenotic papillitis after cholecystectomy, in 16 patients (13.22%) by residual choledocholithiasis, in 12 (9.91%) by choledocholithiasis with chronic calculous cholecystitis and stenosis papillitis, in 6 (4.95%) by choledocholithiasis with stenosis of the terminal choledochitis, in 3 (2.47%) fixedcrement of terminal choledochus and chronic calculous cholecystitis, in 2 (1.65%) - choledocholithiasis with chronic calculous cholecystitis, complicated by cholecystoduodenal fistula, in 5 patients (4.13%) Mirizzi syndrome and in 9 patients (7.43%) exacerbation of pancreatitis on the background of choledocholithiasis with chronic calculous cholecystitis.

The duration of EHC with jaundice was determined according to the E. V. Smirnov classification (1974). Acute cholestasis (up to 10 days) was observed in 35.5% of patients (n=43), prolonged cholestasis (from 10 to 30 days, which after a severe attack took a stable character and lasted up to 4 weeks) was observed in 44.62% of patients (n=54), and chronic (more than 30 days) - in 19.83% (in 24 patients).

To solve the tasks, depending on the type of EHC of non-tumoral genesis of patients was divided into 4 groups: EHC type 1 - without jaundice and without damage to hepatocytes (n=50); type 2 EHC - no jaundice with hepatocyte damage (n=38); type 3 EHC - with jaundice without damage to hepatocytes (n=17); type 4 EHC with jaundice and hepatocyte damage (n=16).

The study excluded patients who had concomitant diseases such as viral and autoimmune hepatitis, Carroll's disease, Wilson-Konovalov's disease, Gilbert's syndrome, and oncological genesis of jaundice. The mean age of the examined patients was 58.23±1.69 years. The youngest patient was 27 years old, the oldest was 83. The largest number of patients were patients aged 40 to 69 years, ie most patients were of working age. Most patients were women (87 (71.9%)).

All patients with simultaneous shear-wave elastometry (SWE) on a Soneus P7 apparatus (Kharkiv, Ukraine-Switzerland) with a sensor with a frequency of 2-5 MHz at a depth of not more than 70 mm from the capsule were performed ultrasound. 7 successful measurements (Σ up Vol. 25, №4, Page 36-43
Structural and functional state of the liver in patients with extrahepatic cholestasis of non-tumor genesis

to 10%, deviation up to 1%) were evaluated, followed by determination of the median, which characterized the stiffness of the liver tissue in kilopascals (kPa). Stratification of reliable results was performed depending on the ratio of IQR/median - less than 30%.

The concentration of total bilirubin in the serum was determined according to the instructions to the kits of EliTech (France). The activity of alanine-aminotransferase (ALT), aspartate-aminotransferase (AST), alkaline phosphatase (ALP), γ-glutamyltransferase (GGT) serum was determined by ultraviolet kinetics (ALT/GPT, AST - AST/ GOT), recommended by the International Federation of Clinical Chemistry (IFCC) according to the instructions of the EliTech kits (France). The presence of endogenous intoxication (EI) was determined by the content of medium molecular weight peptides (MMP) according to V.V. Nikolaychuk. The MMP fraction consists of aromatic amino acids, which are part of proteins, collagen fibers, aromatic amino acids, among which tyrosine and tryptophan occupy a significant place, and therefore the increase in the content of MMP in serum is a marker of activation of catabolic processes in the body. Fibrosis processes were evaluated by the content of hydroxyproline free (HPf) and glycosaminoglycans (GAG). In serum, the content of GAG was determined according to Rimington, HPf - by Osadchuk, the activation of the inflammatory process in patients was evidenced by a change in the level of alpha-1-acid glycoprotein, the content of which was determined by Weimer [16]. Assessment of biochemical parameters was given according to their content in blood of relatively healthy 20 persons (control group).

In order to optimize the mathematical processing, all the input data was entered into a database built using Microsoft Excel spreadsheets. Statistical processing of the obtained data was performed using SPSS 16.0. For statistical analysis of the data used: M - mean value, m - error in determining the mean, comparison of mean values of the variables was performed using parametric method (Student’s t-test) for the normal distribution of these features, expressed in the interval scale. The correspondence of the type of distribution of signs of the law of normal distribution was checked using the Shapito-Wilk method. In other cases, a nonparametric method (the Mana-Whitney U test) was used. The difference in mean values was considered significant at p<0.05. To determine the relationship between the data, a correlation analysis was performed with the calculation of the Spearman correlation coefficient (r) [19].

Results

According to shear wave elastography in patients with EHC of non-tumorogenesis, the stiffness of the liver varied depending on the increase in the intensity of jaundice and was in patients of group I - 6.02±0.223 kPa, II - 6.38±0.171 kPa, III - 7.81±0.321 kPa, in IV - 8.02±0.364 kPa (p<0.001), respectively.

It was found that liver stiffness indices for SWE in patients with EHC differed significantly between the study groups and depended on the level of bilirubinemia (r=0.84) (Fig. 1). The indicators of liver stiffness according to the SWE and the diameter of the choledoch according to the ultrasound are shown in Table 1.

It was found that the liver stiffness indices correlated well with the established choledoch diameter parameters during ultrasound (r = 0.69): I group - 7.58±0.121 mm; II - 8.11±0.191 mm; III - 9.31±0.242 mm; IV - 13.06±0.72 mm (see Table 1 and Figure 2).

Serum bilirubin levels were found to be significantly different in patients with EHC of non-tumorigenic genesis depending on the available jaundice and hepatocyte damage rates, both compared to controls and between the study groups (Fig. 3). Thus, in group II the bilirubin level was 1.5 times higher (34.71±3.23 µmol/l), in III - 6.7 times (159.0±12.7 µmol/l), in IV - 13 times (313.8±28.1 µmol/l) relative to patients of group I (p<0.05).

The activity of serum enzymes confirmed cytolytic processes in the liver. Hyperalanine aminotransferasemia and hyperaspartate aminotransferasemia were observed in patients in all groups compared to controls, but changes

Fig. 1. Dependence of SWE level (kPa) on total serum bilirubin content in patients with non-tumorigenic EHC.

Fig. 2. Dependence of SWE level (kPa) on mean choledoch diameter (mm) in patients with EHC of non-tumorigenic genesis.
in the II and IV groups were more pronounced and probably significant (p<0.001) (Fig. 4).

The determined values of alanine transaminase in serum by groups: 38.22±3.45 U/L in group I; 136.0±27.8 U/L in group II; 39.82±5.11 in group III and 287.0±44.6 U/L in group IV (p<0.001). Serum blood alanine transaminase activity was 36.93±4.14 U/L in group I; 84.13±15.50 U/L in group II; 40.41±6.80 in group III, and 242.0±49.7 U/L in group IV (p<0.001). Such indicators testify to disturbance of permeability of membranes of hepatocytes and their destruction.

Serum cholestasis markers, in contrast to cytolysis indices, had more pronounced pathological changes in patients with non-tumorigenic EHC, both in comparison with the control group and in patients with group IV: their activity increased - alkaline phosphatase to 284.0±34.6 U/L in group I; 262.0±32.8 U/L in group II; 310.0±59.1 U/L in group III and 630.0±81.2 U/L in group IV, respectively, compared with patients in group I (p<0.001) and γ-glutamyltransferase up to 247.0±34.6 U/L in group I; 345.0±37.6 U/L - in group II; 260.0±41.4 U/L in group III; to 610.0±69.9 U/L in group IV, respectively, compared with patients in group I (p<0.001) (Fig. 5).

Patients with EHC of non-tumorigenic genesis had activity of inflammatory processes according to the content of alpha-1-acid glycoprotein in blood serum, which was increased 1.9 times to 0.451±0.041 g/l, (p<0.001) in patients of group I; 2.6 times to 0.632±0.053 g/l - group II;
3.3 times to 0.801±0.053 g/l - group III; 3.8 times to 0.902±0.053 g/l - group IV (p<0.001), respectively, more pronounced pathological changes were characteristic of patients of III and IV groups compared with patients of group I and control indicators, these changes were accompanied by an increase of ESR in patients of group III to 26.70±4.80 mm/g and group IV up to 31.20±3.60 mm/g (p<0.05), as well as an increase in leukocyte levels, which reflected the general inflammatory response of the body (Fig. 6).

The analysis of the data obtained showed a gradual increase in the content of medium-molecular peptides in the serum of patients, depending on the intensity of EHC. The presence of endotoxemia was present in patients of all groups: significant changes were observed in patients of group II, which showed an increase in the concentration of medium-molecular peptides in the serum in 1.3 times to 902.0±56.4 g/l (p<0.01) relative to patients of group I (Fig. 7).

It should be emphasized that in patients of group I, in which the patency disorders were associated with long-term phenomena of EHC without jaundice and lesions of hepatocytes - endogenous intoxication syndrome was formed already at the initial stage of EHC development, and with the increase in bilirubinemia, the content of medium molecular peptides, accordingly, in groups III and IV almost: 1.5 to 1050±76 g/l (p<0.001) and 1.7 times to 1139±78 g/l (p<0.01), respectively, relative to patients of group I; 2.4 and 2.6 times relative to the control group (p<0.001) (see Fig. 7).

There was a gradual decrease in the serum hydroxyproline free content of patients in all groups compared to the control group (Fig. 8). It is impossible to exclude the fact that with the progression of non-neoplastic EHC, there is an imbalance between the synthesis and disintegration of collagen in these patients, which indicates a decrease in the process of collagen degradation, which was more pronounced in patients of group IV, where its content decreased in 2-times to 5,232±0,631 μmol/l (p<0.01) relative to group I (10.72±1.90 μmol/l) and in 1.8 times relative to group II (9.633±1.201 μmol/l, p<0.01) (see Fig. 8).

Increase in GAG concentration in serum was observed in patients in all groups. In group II, this indicator increased 1.4 times to 6.430±0.671 mmol/l (p<0.05); in group III - 1.6 times to 7.133±0.711 mmol/l (p<0.05); in group IV - 1.8 times to 8.140±0.482 mmol/l (p<0.001), respectively, in comparison with patients of group I, indicating the increased breakdown of carbohydrate-protein components of connective tissue, increasing their content in the serum of group II, which showed an increase in the concentration of medium-molecular peptides in the serum in 1.3 times to 902.0±56.4 g/l (p<0.01) relative to patients of group I (Fig. 7).

It should be emphasized that in patients of group I, in which the patency disorders were associated with long-term phenomena of EHC without jaundice and lesions of hepatocytes - endogenous intoxication syndrome was formed already at the initial stage of EHC development, and with the increase in bilirubinemia, the content of medium molecular peptides, accordingly, in groups III and IV almost: 1.5 to 1050±76 g/l (p<0.001) and 1.7 times to 1139±78 g/l (p<0.01), respectively, relative to patients of group I; 2.4 and 2.6 times relative to the control group (p<0.001) (see Fig. 7).

There was a gradual decrease in the serum hydroxyproline free content of patients in all groups compared to the control group (Fig. 8). It is impossible to exclude the fact that with the progression of non-neoplastic EHC, there is an imbalance between the synthesis and disintegration of collagen in these patients, which indicates a decrease in the process of collagen degradation, which was more pronounced in patients of group IV, where its content decreased in 2-times to 5,232±0,631 μmol/l (p<0.01) relative to group I (10.72±1.90 μmol/l) and in 1.8 times relative to group II (9.633±1.201 μmol/l, p<0.01) (see Fig. 8).

Increase in GAG concentration in serum was observed in patients in all groups. In group II, this indicator increased 1.4 times to 6.430±0.671 mmol/l (p<0.05); in group III - 1.6 times to 7.133±0.711 mmol/l (p<0.05); in group IV - 1.8 times to 8.140±0.482 mmol/l (p<0.001), respectively, in comparison with patients of group I, indicating the increased breakdown of carbohydrate-protein components of connective tissue, increasing their content in the serum of group II, which showed an increase in the concentration of medium-molecular peptides in the serum in 1.3 times to 902.0±56.4 g/l (p<0.01) relative to patients of group I (Fig. 7).

It should be emphasized that in patients of group I, in which the patency disorders were associated with long-term phenomena of EHC without jaundice and lesions of hepatocytes - endogenous intoxication syndrome was formed already at the initial stage of EHC development, and with the increase in bilirubinemia, the content of medium molecular peptides, accordingly, in groups III and IV almost: 1.5 to 1050±76 g/l (p<0.001) and 1.7 times to 1139±78 g/l (p<0.01), respectively, relative to patients of group I; 2.4 and 2.6 times relative to the control group (p<0.001) (see Fig. 7).

There was a gradual decrease in the serum hydroxyproline free content of patients in all groups compared to the control group (Fig. 8). It is impossible to exclude the fact that with the progression of non-neoplastic EHC, there is an imbalance between the synthesis and disintegration of collagen in these patients, which indicates a decrease in the process of collagen degradation, which was more pronounced in patients of group IV, where its content decreased in 2-times to 5,232±0,631 μmol/l (p<0.01) relative to group I (10.72±1.90 μmol/l) and in 1.8 times relative to group II (9.633±1.201 μmol/l, p<0.01) (see Fig. 8).

Increase in GAG concentration in serum was observed in patients in all groups. In group II, this indicator increased 1.4 times to 6.430±0.671 mmol/l (p<0.05); in group III - 1.6 times to 7.133±0.711 mmol/l (p<0.05); in group IV - 1.8 times to 8.140±0.482 mmol/l (p<0.001), respectively, in comparison with patients of group I, indicating the increased breakdown of carbohydrate-protein components of connective tissue, increasing their content in the serum of group II, which showed an increase in the concentration of medium-molecular peptides in the serum in 1.3 times to 902.0±56.4 g/l (p<0.01) relative to patients of group I (Fig. 7).

It should be emphasized that in patients of group I, in which the patency disorders were associated with long-term phenomena of EHC without jaundice and lesions of hepatocytes - endogenous intoxication syndrome was formed already at the initial stage of EHC development, and with the increase in bilirubinemia, the content of medium molecular peptides, accordingly, in groups III and IV almost: 1.5 to 1050±76 g/l (p<0.001) and 1.7 times to 1139±78 g/l (p<0.01), respectively, relative to patients of group I; 2.4 and 2.6 times relative to the control group (p<0.001) (see Fig. 7).

There was a gradual decrease in the serum hydroxyproline free content of patients in all groups compared to the control group (Fig. 8). It is impossible to exclude the fact that with the progression of non-neoplastic EHC, there is an imbalance between the synthesis and disintegration of collagen in these patients, which indicates a decrease in the process of collagen degradation, which was more pronounced in patients of group IV, where its content decreased in 2-times to 5,232±0,631 μmol/l (p<0.01) relative to group I (10.72±1.90 μmol/l) and in 1.8 times relative to group II (9.633±1.201 μmol/l, p<0.01) (see Fig. 8).
indicates the activity of the inflammatory process, the duration of which leads to the destruction of liver tissues.

Discussion

The results of the study indicate that in patients with EHC significantly disrupt the biochemical composition of blood serum, which is determined before all types of EHC, which are significantly different in terms of choledochal diameter, liver stiffness, severity of hepatic and cellular insufficiency, EI markers and nonspecific fibrosis markers.

According to SWE, EHC, depending on the available jaundice and hepatocyte damage, definitely affects the elasticity of the liver, regardless of the presence of liver fibrosis/cirrhosis, with an increase in choledoch diameter, liver stiffness and their ratio (r = 0.69) in the case of existing EHC processes are associated with worsening of bile flow, inflammatory phenomena in the ductal system and, as a consequence, increased hydrostatic pressure in the ducts and edema of the liver parenchyma, which is confirmed by other studies [11-13, 20-22, 24-27].

The liver is one of the first organs whose functions are altered as a result of impaired bile flow in EHC of non-tumorigenesis. The presence of signs of cholestasis was associated with more significant changes in the activity of serum enzymes of serum, with a more pronounced cytolytic and cholestatic syndrome with increasing intensity of cholestasis. A similar pattern was observed with regard to the content of α1-acid glycoprotein in blood serum, the value of which was highest in patients of group IV with non-neoplastic EHC and showed activation of inflammatory processes in the body, revealed changes were accompanied by an increase in leukocyte and ESR levels inflammatory response of the body.

An important pathogenetic syndrome of homeostasis disorders in virtually all diseases is endogenous intoxication - an integral concept that includes a number of components: the accumulation of intermediates of impaired metabolism and metabolites, endogenous and bacterial toxins, aggressive biologically active substances, inflammatory factors in combination with hypoxia and disorders of microcirculation [14, 15, 23].

Occurrence in the blood of non-oxidized products of lipid metabolism (acetone, acetoacetic and β-Hydroxybutyric acid), increase in the level of ketones, medium-molecular peptides cause the development of the clinical picture of endogenous intoxication syndrome. The obvious important role of the liver, as a body of metabolic detoxification, in the process of formation of EI in the body, as well as the processes of cytolysis and cholestasis.

Therefore, extrahepatic cholestasis of non-tumoral genesis is accompanied by certain metabolic disorders and ultrastructural changes that lead to impaired cell-tissue metabolism, the appearance of a large number of aggressive active substances, the reduction of the inactivation and elimination of toxins, that accumulate in the patient and leads to the development of general endotoxemia.

Given the data obtained, it is promising to further study liver stiffness under conditions of surgical correction as a non-invasive way of monitoring the syndrome of "rapid discharge" of bile in patients with non-hepatic cholestasis of non-tumoral genesis.

Conclusions

1. Indicators of liver stiffness according to SWE data in patients of all groups with extrahepatic cholestasis of non-tumoral genesis depend on the level of bilirubinemia (r = 0.84), and in patients with hepatocyte damage (II and IV group) also on the duration of cholestasis (r = 0.69) but cannot serve as an objective indicator of liver functional status.

2. Patients with extracorporeal cholestasis of non-tumoral genesis without the appearance of jaundice and hepatocyte damage revealed an increase in endotoxia indicators (p<0.001), which indicates developing of EHC.

3. Extrahepatic cholestasis of non-tumorigenesis is accompanied by a gradual increase in alkaline phosphatase and γ-glutamyltransferase activity, while bilirubin levels cannot serve as an indicator of long-term EHC.

References

Гепатоциты.

Согласно статистике, гепатоциты в организме человека являются основными структурными элементами печени. Они ответственны за многие функции, включая производство билирубина, непосредственно связанного с желчью. Билирубин функционирует как естественное индикаторное средство для оценки длительности холестаза. Билирубинное накопление вызывает увядание гепатоцитов, что может привести к их смертельной переработке.

Важно отметить, что билирубин не является независимым индикатором длительности холестаза. Уровень билирубина может быть повышен в результате множества факторов, включая непосредственный повреждение гепатоцитов. Повреждение гепатоцитов может быть обусловлено различными факторами, включая хронический алкоголизм и наследственные заболевания.

Другим важным механизмом является интраэпителиальный холестаз, который может приводить к повреждению гепатоцитов. Этот механизм включает в себя процесс накопления холестерина в гепатоцитах, что приводит к повреждению клеток и развитию холестаза.

Структурно-функциональный статус печени в условиях позапечёночного холестаза.

В таблице представлены данные по структурно-функциональному состоянию печени в условиях позапечёночного холестаза, а также показатели длительности холестаза и выхода из него.

<table>
<thead>
<tr>
<th>Название</th>
<th>Показатели</th>
</tr>
</thead>
<tbody>
<tr>
<td>Позапечённый холестаз</td>
<td>Билирубин (μmol/L)</td>
</tr>
<tr>
<td>Позапечённый холестаз</td>
<td>Суточное потребление билирубина (μmol/L)</td>
</tr>
<tr>
<td>Позапечённый холестаз</td>
<td>Трансаминазы (мкЕ)</td>
</tr>
<tr>
<td>Позапечённый холестаз</td>
<td>Уровень билирубина в сыворотке крови (μmol/L)</td>
</tr>
</tbody>
</table>

Ключевые слова: позапечёночный холестаз, непушенное генез, временные гепатоциты.
СТРУКТУРНО-ФУНКЦИОНАЛЬНОЕ СОСТОЯНИЕ ПЕЧЕНИ У БОЛЬНЫХ С ВНЕПЕЧЕНОЧНЫМ ХОЛЕСТАЗОМ НЕОПУХОЛЕВОГО ГЕНЕЗА
Шевченко Б.Ф., Зеленюк А.В., Кленина И.А., Бабий А.М.
На сегодняшний день диагностика и лечение внепеченочного холестаза (ВПХ) на разных этапах развития остается одной из самых актуальных проблем современной билиарной хирургии. Цель исследования - установить структурные изменения печени у пациентов с внепеченочным холестазом неопухолового генеза по данным сдвиговолновой эластометрии и ультразвукового исследования в сопоставлении с биохимическими показателями эндотоксемии, воспаления и маркерами фиброза. Предоперационно обследовали 121 больного с ВПХ неопухолового генеза провели стандартные общеклинические исследования, ультразвуковое исследование печени и ее сдвиговолновую эластометрию. В зависимости от наличия желтухи и симптомов повреждения гепатоцитов всех исследованных разделены на 4 группы. Определяли содержание в сыворотке крови среднемолекулярных пептидов, свободного гидроксипролина и гликозаминогликанов. Статистическую обработку полученных данных проводили с использованием программы SPSS 16.0. У пациентов с ВПХ неопухолового генеза по данным сдвиговолновой эластографии жесткость печени менялась в зависимости от увеличения интенсивности желтухи. Установлено, что уровень билирубина в сыворотке крови у таких пациентов достоверно отличался в зависимости от имеющейся желтухи и симптомов повреждения гепатоцитов как по сравнению с данными контрольной группы, так и между исследованными группами. При ВПХ структурные изменения печени зависят от нарушений биохимического состава сыворотки крови, которые существенно изменяются при длительном внепеченочном холестазе и проявляются выраженной печеночной недостаточностью. Таким образом, у пациентов с имеющимися ВПХ неопухолового генеза без желтухи и повреждения гепатоцитов выявлено увеличение показателей эндотоксемии (p<0,001), что свидетельствует о развитии холестаза; показатели жесткости печени зависят от уровня билирубинемии (r = 0,84), а у пациентов с повреждением гепатоцитов еще и от длительности холестаза (r = 0,68). Уровень билирубина не может быть индикатором длительности внепеченочного холестаза.

Ключевые слова: внепеченочный холестаз неопухолового генеза, эластометрия, маркеры эндогенной интоксикации, повреждение гепатоцитов.