Ito stellate cells (hepatic stellate cells) in diagnosis of liver fibrosis

Introduction

Adverse outcome of most chronic diffuse liver lesions of various etiologies including chronic hepatitis C (CHC) is liver fibrosis in the development of which main participants are activated fibroblasts. Activated hepatic stellate cells (HSC) are the main source of activated fibroblasts.1,2

HSC is a synonym - stellate cells of liver, Ito cells, perisinusoidal lipocytes, stellate cells. HSC were firstly described in 1876 and named by K. Kupffer stellate cells T. Ito, finding in them a drop of fat firstly marked them as lipophagic (“shibo-sesshusaibo”) but then finding that the fat elaborated by the cells from glycogen named them as fat-storing cells or Ito sells (“oshibo-chozosaibo”).1 In 1971, K. Wake proved the identity of stellate cells of Kupffer and Ito cells and that these cells are “stored” vitamin A.4

About 80% of vitamin A which is in the body accumulates in the liver and up to 80% of liver retinoids deposited in fat droplets of HSC. Ethers of retinol in formulation of chylomicrons entered to hepatocytes, where they are converted to retinol and form of vitamin A complex with retinol binding protein (RBP), which is secreted in perisinusoidal space where it deposited by cells.5

Established by K. Popper close relationship of HSC with the hepatic fibrosis demonstrated theirs not static, but dynamic function. It means the ability to be directly involved in the remodeling of intralobular perhepatocellular matrix.6 The main method of morphological study of liver conducted by assessing the changes in lifetime biopsy materials is light microscopy. This method in clinical practice allows establishing the stage of inflammation activity and stage of chronization.7 The disadvantage of this method is low resolution which does not allow evaluating features of cell structure, intracellular organelles, insertions, functional characteristics. Lifetime electron-microscopic study of ultrastructural changes in the liver makes it possible to supplement the light microscopy data and raise their diagnostic value.

In this regard the identification of the HSC, study of their phenotype during transdifferentiation and determining the intensity of their proliferation are the major contribution to the prediction of outcomes of liver disease as well as in the patomorphology and pathophysiology of fibrogenesis. The aim of research is to introduce structural and functional description of HSC on the results of cytological identification of lifetime liver biopsies.

Abbreviations: CD, cluster of differentiation; CHC, chronic hepatitis C; HCV, hepatitis C virus; HSC, hepatic stellate cells; ICAM, intercellular adhesion molecule; MMPs, matrix metalloproteinases; PDGF, platelet-derived growth factor; RBP, retinol-binding protein; RER, rough endoplasmic reticulum; TIMPs, tissue inhibitors of matrix metalloproteinases

Methods

Lifetime liver biopsy obtained by performing aspiration biopsy of the liver of patients with CHC (HCV ribonucleic acid+). Patients gave theirs written informed consent. For light microscopy semifine sections of liver biopsy samples of patients 0,5x2mm in size were fixed by double fixation method: firstly by the Sato Taizan technique.8 Then tissue samples for 1 hour additionally fixed in 1% osmium fixative prepared with 0.1M phosphate buffer of Zerensen, pH 7.4. For better identification of intracellular structures and interstitial substance on semifine sections in 1% osmium tetroxide was added potassium dichromate (K2Cr2O7) or crystals of chromic anhydride (1mg/ml). After dehydration of samples in series of alcoholic solutions of increasing concentrations and acetone they were placed in the prepolymerized mixture of butyl methacrylate and styrole and polymerizing at 0,5°C. For further study of ultrastructural changes. Ultrathin sections (35 nm) were prepared semifine sections. Ultrathin sections (400 nm) and stained with methylene blue. Electron-microscopic spesimens were embedded in Araldite from these blocks on ultramicrotome Leica EM VC7 (Germany) were prepared semifine sections (400 nm) and stained with mylene blue. Specimens viewed under the light microscope and the same type of slide chosen for further study of ultrastructural changes. Ultrathin sections (35 nm) contrasted with 2% solution of uranyl acetate per 50% of methanol and lead citrate by E.S. Reynolds.11 Electron-microscopic spesimens were studied in the electron microscope JEM-1011 (JEOL, Japan) with an increase of 10000–60000 at an accelerating voltage of 80 kV.
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For images was used a set of digital camera Olympus MegaViewIII (Germany) and programs for processing of images iTEM (Olympus, Germany).

Results and discussion

HSC located in perisinusoidal space (Disse) in recesses between hepatocytes and endotheliocytes and have long processes that penetrate deep between hepatocytes. In the majority of publications devoted to this population of HSC is a schematic representation of them which allows only indication of “territorial” belonging of HSC in the liver and in relation to others of their “neighbors” (Figure 1).

Figure 1 Diagram of HSC location in perisinusoidal space of Disse, an internete resource.

HSC have close contact with endotheliocytes through the incomplete basement membrane components and interstitial collagen fibers. Nerve endings penetrate between HSC and parenchymatous cells, wherefore the space of Disse defines as the space between the plates of parenchymatous cells and HSC complex and endotheliocytes.14

It is supposed that HSC become from poorly differentiated mesenchymal cells transverse septum of developing liver.15 The experiment revealed that in the formation of HSC hematopoietic stem cells participates, and that this process is not caused by cell fusion.16

Sinusoidal cells, first of all HSC, play a leading role in all forms of liver regeneration. Fibrotic liver regeneration occurs as a result of the inhibition of HSC stem functions and stem cells of marrow.17 In human liver HSC constitute 5-15% being one of 4 types of sinusoidal cells with mesenchymal origin: Kupffer cells, endotheliocytes and Pit-cells. Within pool of sinusoidal cells 20-25% of leukocytes are detected.18,19

There are lipid inclusions with retinol, triglycerides, phospholipids, cholesterol, free fatty acids, a-actin and desmin in the cytoplasm of HSC.20 To visualize HSC auric chloride staining is used. The experiment found that differentiation marker of HSC from other myofibroblasts is the expression by them protein reelin.21

HSC exist in a stable - “inactive HSC”, transitional and long-term active condition, each of which is characterized by genes expression and the phenotype (alpha-smooth muscle actin, intercellular adhesion molecule (ICAM)-1, chemokines and cytokines).

HSC in stable condition are round, slightly elongated or irregular shape, large nucleus and vivid imaging feature - lipid inclusions (droplets) containing retinol (Figure 2).

The amount of lipid droplets in the inactive HSC is 30 or more.

They are similar in size and are adjacent to each other, pressing into the nucleus and pushing it aside periphery (Figure 2). Small inclusions can be placed between large droplets. The color of droplets depends on the fixator and staining of material. In one case, they are bright (Figure 2A), in the other they are dark green (Figure 2B).

At electron microscopy on a background of the light lipidic substance more osmiophil marginal ring is formed (Figure 2C).22 Most of the “inactive” HSC along with large lipid inclusions small amount of cytoplasmic matrix poor on mitochondria and rough endoplasmic reticulum (RER) is seen. However, compartments of moderately developed Golgi complex in the form of stacks of 3-4 flattened cisterns with slightly extended ends are clearly seen (Figure 2S).

Figure 2 HSC which are in inactive condition.

(A) Hepatic stellate cells (HSC) of round shape with a high content of lipid droplets with a light color (white arrows), hepatocytes (Hep) with the devastated cytoplasm (black arrow); B, HSC with lipid droplets of dark color (white arrows), in close contact with macrophage (M); A, B, Semfine sections, stained with azure II - basic fuch sine, microphotography, zoom 1000; C, HSC with plenty of lipid droplets (more then 30) having an irregular shape, zoom 6000 (Hep – hepatocyte); D, Ultrastructural components of HSC: lipid droplets (L), mitochondria (orange arrow), rough endoplasmic reticulum (green arrows), the Golgi complex (red arrow), zoom 15000; (C-D) Electron diffraction patterns.

Under certain conditions activating HSC obtain mixed or transitional phenotype combining morphological characteristics of lipid like and fibroblast-like cells (Figure 3).22

The transitional phenotype of HSC also has its morphological features. The cell becomes elongated shape, the number of lipid inclusions decreases and there is a reduction of intussusception of the volume of the cytoplasm containing numerous cisterns of RER with related ribosomes and free ribosomes and mitochondria is increasing. There is hyperplasia of lamellar Golgi complex components’ represented by multiple stacks of 3-8 flattened cisterns and the number of lysosomes involved in degradation of lipid droplets is increasing (Fig. 3D). Hyperplasia components of RER and Golgi complex associated with ability of fibroblasts to synthesise

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collagen molecules, as well as to simulate them by post-translational hydroxylation and glycosylation in the endoplasmic reticulum and Golgi complex elements.

Figure 3 HSC are there in the transitional state.

Hepatic stellate cells (HSC) (white arrows), semifine sections, stained with azure II - basic fuchsin, microphotography, zoom 1000; B, HSC of elongated shape and a small amount of lipid droplets, zoom 8000 (Hep – hepatocyte, RBC – red blood cell, LYM – lymphocyte); C, HSC in contact with Kupffer cells (KC) and lymphocyte (LYM), zoom 6 000 (Hep – hepatocyte); D, Mitochondria (orange arrow), rough endoplasmic reticulum (green arrows), the Golgi complex (red arrow), lysosomes (blue arrows), zoom 20 000; B, C, D, Electron diffraction patterns.

In the intact liver HSC while at rest cover by their processes sinusoidal capillary. Processes of HSC divided into 2 types: perisinusoidal (subendothelial) and interhepatocellular (Figure 4).

Perisinusoidal (subendothelial) processes out of the cells’ body and extend along the surface of sinusoidal capillary covering it by thin digitiform rami. They are covered with short villi and have characteristic long micro-emissions extending even further on the surface of endothelial capillary tube. Interhepatocellular outgrowths, breaking the lamella of hepatocytes and attaining neighboring sinusoid are divided into several perisinusoidal outgrowths. Thus, HSC on average cover over two adjacent sinusoids.

When liver damages then activation of HSC and fibrogenesis occur which distinguish 3 phases. They are designated as the initiation, prolongation and the resolution (resolution of fibrous tissue).

This process of transformation of the “stable” HSC in fibrosing myofibroblasts is activated by cytokines (interleukin-1, interleukin-6, tumor necrosis factor alpha), insufficiently oxidized products of metabolism, active form of oxygen, nitric oxide, endothelin, platelet-derived growth factor (PDGF), plasminogen activator, transforming growth factor β-1, acetaldehyde and others. Direct activators are hepatocytes in state of oxidative stress, Kupffer cells, endotheliocytes, leukocytes, platelets which produce cytokines (paracrine signals) and HSC themselves (autocrine stimulation). The activation is accompanied by the expression (inclusion in work) of new genes, the synthesis of cytokines and extracellular matrix proteins (collagen I, III, V type).

At this stage the process of HSC activation can be completed due to the stimulation of formation in HSC antiinflammatory cytokines inhibiting tumor necrosis factor alpha production by macrophages in the damage zone. As a result, the amount of the HSC is sharply reduced, they undergo apoptosis and processes of fibrosis in the liver do not develop.

At the second phase (prolongated) during prolonged continuous paracrine and autocrine influence of activating stimuli in the HSC “supported” an activated phenotype characterized by the transformation of HSC in contractile myofibroblast-like cells which realize synthesis of extracellular fibrillar collagen.

The activated phenotype is characterized by proliferation, chemotaxis, contractility, loss of retinoid reserves and formation of cells which are myofibroblast-like. Activated HSC also demonstrate high content of new genes such as alpha-smooth muscle actin, ICAM-1, chemokines and cytokines. Cell activation indicates the beginning of the early stage of fibrogenesis and precedes increased production of extracellular matrix proteins. Formed fibrous tissues undergo remodeling due to the splitting of the matrix with help of matrix...
metalproteinases (MMPs). In turn, splitting of matrix is regulated by tissue inhibitors of MMPs (TIMPs). MMPs and TIMPs are part of a family of zinc-dependent enzymes. MMPs are synthesized as inactive HSC proenzymes which are activated by removal of propeptide, but inhibited by interaction with endogenous TIMPs - TIMPs-1 and TIMPs-2. HSC produce 4 types of membrane-type MMPs which are activated by influence interleukin-1. Among MMPs particular importance is attached to MMPs-9 - neutral matrixmetalloproteinase which has activity against collagen of type 4, which is the part of basal membrane, as well as against partially denatured collagens 1st and 5th types.30-32

About the increase in population of HSC in various liver damage is judged by an activity of significant number of mitogenic factors, related to them tyrosine kinase receptors and other identified mitogens, which cause the most pronounced proliferation of HSC: endothelin-1, thrombin, fibroblast growth factor, PDGF, insulin-like growth factor. The accumulation of HSC in areas of damaged liver occurs not only due to the proliferation of these cells, but also due to their directional migration to these areas by chemotaxis, with the participation of such chemoattractants, as PDGF and leukocyte chemoattractant – monocyte chemotactic protein-1.24,32

At activated HSC number of lipid droplets is reduced to 1-3 with their location on opposite poles of the cell (Figure 5).

Figure 6 High fibrotic activity of HSC.
(A-B) Myofibroblasts (MFB), the cell contains large nucleus, elements of rough endoplasmic reticulum (red arrows), numerous free ribosomes, polymorphic vesicles and granules, single mitochondria and bright imaging feature - a bundle of actin filaments in cytoplasm (yellow arrows), zoom 12 000 and 40 000 (Hep – hepatocyte). (C-F) High fibrotic activity of Hepatic stellate cells (HSC) while maintaining at cytoplasm retinoid containing lipid droplets. Numerous bundles of collagen fibrils (white arrows), save (C) and lost (D-F) specific cross striation; zoom 25 000, 15 000, 8 000, 15 000, electron diffraction patterns.

In 2007 for the first time HSC were named as liver stem cells since they express one of the markers from a hematopoietic mesenchymal stem cells - cluster of differentiation (CD)-133.31

Myofibroblasts which are absent in normal liver have three potential sources: the first - at intratramer liver development in portal tracts myofibroblasts surround blood vessels and bile ducts during their maturation and after the full development of the liver they disappear and are replaced in the portal tracts by portal fibroblasts; the second - with the damaging of liver they are formed due to the portal mesenchymal cells and resting HSC. Sometimes they are formed due to expense of the transitional epithelial-mesenchymal cells. They are characterized by the presence of CD45-, CD34-, Desmin+, glial fibrillary acidic protein+ and Thy-1+.

Recent studies have shown hepatocytes, cholangiocytes, and endothelial cells can become myofibroblast through epithelial- or endothelial-mesenchymal transition. These cells include such markers as CD45-, albumin + (i.e. hepatocytes), CD45-, CD19+ (i.e. cholangiocytes) or Tie2+ (endothelial cells). In addition, marrow cells consisting of fibrocytes and circulating mesenchymal cells may transform into myofibroblasts. These cells are CD45 + (fibrocyte), CD45 +/- (circulating mesenchymal cells), collagen type I+, CD11d+ and major histocompatibility complex, class II+ (Figure 7).2,15

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Figure 7 Participants of miofibroblastic activation of HSC.

(A) Interaction of ductal hepatic precursor cells with myofibroblasts of liver; (B) Miofibroblast-like Hepatic stellate cells (HSC) in condition of fibrogenic activation.34,35

Literature data confirm not only the close relationship proliferation of oval cells with proliferation of sinusoidal cells,36,37 but also the data about possible differentiation of HSC in the hepatic epithelium36 which was called epithelial-mesenchymal transformation of perisinusoidal cells.37

In the state of fibrogenic activation miofibroblast-like HSC along with decrease the number and following disappearance of lipid droplets characterized by nodal proliferation (Figure 8), the immunohistochemical expression of fibroblast-like markers including smooth muscle α-actin and formation of pericellular collagen fibrils in the space of Disse.34,35

Figure 8 Nodal proliferation of HSC in condition of fibrogenic activation.

(A) Hepatic stellate cells (HSC) hyperplasia (white arrows) in the lumen of dilated sinusoids (Hep – hepatocyte); (B) Proliferation of transdifferentiational HSC (white arrows), endothelial cell (pink arrow). (A,B) Semifine sections, stained with azure II - basic fuchsine, microphotography, zoom 1000.

In phase of fibrosis increasing hypoxia of liver tissue becomes a factor of additional surplus expression in stem cells of pro-inflammatory adhesion molecules - ICAM-1, ICAM-2, vascular endothelial growth factor, pro-inflammatory chemoattractants - monocyte colony stimulating factor, monocyte chemotactic protein-1 and cytokine-induced neutrophil chemoattractant and others that stimulate the formation of pro-inflammatory cytokines (transforming growth factor β-1, PDGF, fibroblast growth factor, PAF, SCF, ET-1) and strengthen the processes of fibrogenesis in the liver creating the conditions for a self-supporting induction, ongoing activation of HSC and process of fibrogenesis.32

On microscopic specimens pericapillary fibrosis displays as intense color of perisinusoidal connective tissue and layer of intercellular matrix surrounding hepatocytes (they are often die) in red. At the electron microscopic specimens fibrotic changes visualized either in the form of formed large fascicles of fibrils of collagen fibers preserved diametrical bending, or in the form of massive deposits in the Disse space of fiber mass, which is a swollen and lost periodic banding of collagen fibers (Figure 9).

Figure 9 The final stage of miofibroblastic activation of HSC.

(A-B) Perisinusoidal fibrosis (white arrows). Perisinusoidal connective tissue and layer of intracellular matrix surrounding hepatocytes (Hep) stained with basic fuchsine in red. Hepatic stellate cells (HSC) activated and transformed in fibroblasts (blue arrows). Hepatocyte at picture A – hepatocyte with devastated cytoplasm. Semifine sections, stained with azure II - basic fuchsine, microphotography, zoom 1000.

(C-D) Perisinusoidal and perihepatocellular fibrosis in the liver lobule, increased electron density of fibril collagen fibers; condensation of matrix of mitochondria in hepatocyte (orange arrow). Zoom 8 000 and 15 000 respectively, electron diffraction patterns.

On microscopic specimens pericapillary fibrosis displays as intense color of perisinusoidal connective tissue and layer of intercellular matrix surrounding hepatocytes (they are often die) in red. At the electron microscopic specimens fibrotic changes visualized either in the form of formed large fascicles of fibrils of collagen fibers preserved diametrical bending, or in the form of massive deposits in the Disse space of fiber mass, which is a swollen and lost periodic banding of collagen fibers (Figure 9).
According to modern concepts fibrosis is a dynamic process that can progress and regress (Figure 10). Recently, several specific markers of HSC have been proposed: the rise of vitamin A in lipid droplets, glial fibrillary acidic protein, nerve growth factor receptor and synaptophysin.25,41 Researches on participation of liver HSC in proliferation and differentiation of stem cells of the liver are taken.13,16 We studied level of retinol and the content of RBP4 at blood serum. RBP4 forms a complex with retinol and whose concentration in blood plasma correlates normally with the body’s provision of vitamin A, 80% of which is in the HSC (Table 1).

![Image](51x515 to 284x631)

**Figure 10** Dynamic of events in the pseudolobule of patient with viral cirrhosis of liver 6 months after transplantation in liver autologous mesenchymal stem cells. (A) Pseudolobules surrounded by fibrous septa, staining by Masson, zoom 50; (B) Resolution of fibrous septa around pseudolobules, staining by Masson, zoom 200.

| Group               | n  | RBP4, ng/ml | Vitamin A, mcmole/l |
|---------------------|----|-------------|---------------------|
|                     |    | M± m        | p                   |
| Liver cirrhosis     | 17 | 23.6±2.29   | <0.05               |
| CH, AST norm        | 16 | 36.9±2.05*  | >0.05               |
| CH, AST >2 norm     | 13 | 33.0±3.04*  | >0.05               |
| CH, ALT norm        | 13 | 37.5±3.02*  | >0.05               |
| CH, ALT >2 norm     | 21 | 35.9±2.25*  | >0.05               |
| Control             | 15 | 31.2±2.82   | 1.22±0.077          |

Note: p - significant differences with the control (p<0.05); * - Significant differences between liver cirrhosis and chronic hepatitis (CH) (p<0.05)

Dependence between the content of RBP4 and retinol with the 4-th stage of fibrosis (cirrhosis) is fixed. In contrast to chronic hepatitis dependence between the content of RBP4 and 4-th stage is not traced, regardless of the biochemical markers of inflammatory activity in liver.

We think that the development of hyper-retinemia in liver cirrhosis is due to fibrotic evolution and sharp decrease in volume of parenchymal component of liver. And as a result of which the retinol is “washed out” of the HSC because of their degranulation and accumulation of its high level in blood content. This process is associated with a decrease in protein-synthesizing function of liver, which leads to the decrease in the synthesis of RBP4 and its inadequate accumulation in the liver. As a consequence, the deficiency of retinol in liver contributes to the progression of fibrosis.

### Conclusions

Maximum effectiveness of structural and functional evaluation of HSC provides by morphological study of intravitral biopsy with simultaneous use of complex cellular imaging techniques (light and electron microscopy of ultrathin sections and original methods of fixation and staining). Results of morphological studies of HSC will improve the quality of intravitral diagnosis of fibrosis, to hold its monitoring and prediction of outcomes chronic diffuse liver lesions at a higher level today. Results of morphological findings will help the clinician furtherly include in wording of final diagnosis more accurate data on the chronic stage (stabilization, progression or resolution of fibrosis) in the course of therapy.

### Author contributions

a. V Tsyrkunov: study concept and design, performed experiments, collected data, analyzed and interpreted data, wrote all drafts of the manuscript, and revised the manuscript, study supervision.

b. V Andreev: performed experiments, collected data, wrote all drafts of the manuscript.

c. R Kravchuk: performed experiments, collected data, wrote all drafts of the manuscript.

d. I Kandratovich: performed experiments, collected data, performed statistical analysis, and wrote all drafts of the manuscript.

### Conflicts of interest

The authors have no conflicts of interests related to this publication.

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