Update of Key Mechanism of Hepatic Peliosis

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ABSTRACT

Hepatic peliosis is a rare vascular histological feature characterized by dilatation and rupture of sinusoidal blood-filled spaces within the liver. There may be occur on other organs. It observed in human and occurred in aged rat. There is a complex interplay could share in peliosis formation. Loss of HGF/c-met mediate fas apoptosis which may be a part of axis explain peliosis hepatica. Additional knockdown of hepatocyte growth factor widening sinusoidal space and enhanced hepatocyte dissociation. Notably, Egfr and Tnfr1 also mediate fas apoptosis and could be a part of key mechanism of peliosis. Also peliosis seen in transgenic model like Hgf transgenic model treated with diethyl nitrosamine and c-myc hepatocellular carcinoma of c-myc transgenic model of liver cancer; while vegf have dual role in protection or induction of liver peliosis. Taken all, many factors could play a role in induction of liver peliosis.

Keywords: Hepatic Peliosis; HGF/c-Met; Fas; Vegf

Introduction

Hepatic peliosis is a rare vascular condition characterized by dilatation and rupture of sinusoidal blood-filled spaces within the liver. There may be occur on other organs like spleen, bone marrow, lungs, and abdominal lymph nodes Downes et al. [1], peliosis identified by computed tomography of the abdomen of human case revealed a large well-defined lesion with heterogeneous density in the right lobe of the liver compatible with intrahepatic hemorrhage in the absence of definite extravasation (Charatcharoenwitthaya and Tanwandee, 2014). Also, spontaneous hepatic rupture, occur during pregnancy in a patient with peliosis hepatis Cimbanassi et al. [2]. Peliosis hepatis, another reactive vascular lesions which radiologically similar to liver tumors, is characterized by focal or diffuse cystic blood-filled cavities mainly localized in parenchyma. Peliotic lesions show a loss of endothelial lining and replaced by perisinusoidal reticulin network. Peliosis hepatis can be caused by a wide array of factors, including chemicals, infections and drugs as recently recorded that Hepatorenal peliosis a characteristic feature of gentamox antibiotic induce toxicity in albino rats Elalfy et al. [3] and also the corticosteroid drugs induced peliosis hepatica in Systemic Lupus Erythematosus Zimmermann [4] Kimura et al. [5].

Hepatic peliosis named spongiosis hepatis is a spontaneously occurring lesion in the livers of ageing rats, appearing most often in the second year of life, with a strong predilection for male animals. The incidence of spontaneous spongiosis hepatis can reach 34% in male Fischer rats. The lesion is composed of altered sinusoidal lining cells which some authors have interpreted as hepatic stellate cells (Ito cells or fat-storing cells). The characteristic histological appearance is described by Bannasch et al. [6-8] in rats treated with carcinogens - spontaneous spongiosis hepatis is less likely to be multifocal and is less often associated with stellate cell aggregates. In contrast, electron microscope scanning revealed that hepatic stellate cells become enlarged and show apoptosis in space of dis under effect of gadolinium chloride and diethylnitrosamine induced hepatocellular carcinoma treated balb-c mice El-hadidy et al. [9]. So may be enlarged HSC responsible of widening sinusoidal space, collection of blood into small cavities and share in peliosis mechanism.

Also, Elalfy et al. [10] found that egf hepatocellular carcinoma show peliosis like appearance pathological description of peliosis by light microscopy as a centrilobular glycogen depletion and cytoplasmic foaminess, usually followed by extensive vacuolation and centrilobular congestion and hydropic single cell necrosis on mice treated with furosemide (walker). While by Electron microscopy revealed disaggregation of polyribosomes, vesiculation...
of endoplasmic reticulum, an endocytic origin for the vacuolation, and a definite sequence in the development of congestion. Walker et al. [11] the gross inspection of the peliotic lesions show the cut sections a “swiss cheese” appearance. Microscopically, there were two different types of peliosis can be identified in the liver: first type, parenchymal peliosis, composed of irregular cavities that outer lining was not be sinusoidal cells or fibrous tissue; and the second one, phlebitic peliosis, was regular, spherical cavities protected by endothelium or fibrosis Tsokos et al. [12]. The causes of peliosis hepatica is unknown and it occasionally observed in human liver Craig et al. [13] Many author interests to better understand explanation of liver peliosis as Munoz et al. [14] explain Peliosis Hepatitis to super expression of fas legend a Complication of the Ise of Oral Contraceptives in a Patient with Myelodyplasia.

Knockdown of endothelial Fas sufficiently recapitulated the protection against hemorrhage seen with the addition of mural cells. the regulation of endothelial Fas signaling is involved in the promotion of vascular integrity by mural cells in tumors Kamei et al. [15]. Moreover, HGF is a potent angiogenic factor in vitro and in vivo Bussolino et al. [16,17], and is involved in haematopoiesis Zarnegar et al. [18] and local regulation of fibrinolysis and coagulation Pepper et al. [19] Wojta et al. [20]. Some of HCCs in Hepatocyte growth factor transgenic mice treated with diethyl nitrosamine were accompanied peliosis-like change Horiguchi et al. [21]. The most consistent abnormality of HGF knockout mice was a loosened liver structure with enlarged sinusoidal spaces and dissociation of the parenchymal cells. The dissociated cells often showed signs of apoptosis Shiota et al. [22]. Notably importance of TGF-beta signaling in the control of liver homeostasis and there was no peliosis seen in histology feature of c-Met (cyo-Met) and c-Myc transgenic model Amicone et al. [23]. The HCCs observed in c-myc transgenic mice were either of the trabecular or of the solid histological type, varying from well differentiated to poorly differentiated tumors with cell polymorphism, atypia and areas of hemorrhagic necrosis Thorgeirsson et al. [24,25], moreover, c-myc transgenic mice enhanced loss of c-met receptor for hepatocyte growth factor ligand after one year aged mice Thorgeirsson et al. [24,25] C-Met activation by HGF impairs Fas-triggered apoptosis of primary embryonic hepatocytes and cell survival correlates with inhibition of caspase-8 and caspase-3 activities. HGF treatment prevents degradation of FLIPL triggered by Fas activation.

In contrast to this, c-Met activation does not modulate FLIPL levels and its stability in untreated cells Moumen et al. [2007]. Also, c-Met, a growth factor receptor tyrosine kinase a cell survival mechanism, directly binds to and sequesters the death receptor Fas in hepatocytes. This interaction prevents Fas self-aggregation and Fas ligand binding, thus inhibiting Fas activation and apoptosis Wong et al. [26]. Loss of c-Met function increased sensitivity to Fas-mediated apoptosis Gómez Quiroz et al. [27]. Fas and loss of hepatocyte growth factor, a potent hematopoietic agent, may be among the factor enhanced peliosis occurrence and complex interplay of pro-apoptotic signals and have shown to be Fas mediated Feldmann et al. [28]. Specifically, loss of hepatocyte growth factor receptor signaling sensitizes to Fas-mediated apoptosis Gómez Quiroz et al. [27]. As Fas was strongly induced in HCC of transgenic mice that had peliosis feature histological, at least in part, by Fas mediated apoptosis of hepatic cells or epidermal growth factor receptor mediated CD95 tyrosine phosphorylation for formation of death disc, the death-inducing signaling complex Reinehr et al. [29,30]. Furthermore, EGFR ligands and TNF receptor 1 sensitizing hepatocyte apoptosis during fulminate hepatitis and for TNF signaling to mediated Fas –apoptosis Murthy et al. [31].

Finally, c-met regulating gene, Ets 1,2 are known regulate endothelial function and gene expression. For example, Ets1 has been shown to modulate the expression of several genes of endothelial function and angiogenesis including VEGF-R2, VEGF, and Tie2.2 However, knockdown of Ets1 is associated with defects in T-cell function, but no abnormalities in vascular development or angiogenesis. The protective agent against peliosis was recorded in rat intoxication with cadmium chloride Tzirogiannis et al. [32] by Putrescine or VEGF administration totally reversed macroscopic peliosis. Putrescine considered a major protective effect on hepatocytes, whereas the protective effect of VEGF was more significant for nonparenchymal liver cells Tzirogiannis et al. [33]. Additionally, The hepatoprotective effect of Hepatic Stimulator Substance against cadmium-induced necrosis, apoptosis, and peliosis Tzirogiannis et al. [34]. Additionally. Gene knockdown experiments have stated the potential roles played by Vascular Endothelial Growth Factor (VEGF) VEGF receptor 2 (VEGF-R2) and VEGF-R1 in new vessel formation Biscetti et al. [35-37] and have reported that PPAR alpha and PPAR gamma activation enhanced neo-angiogenesis through (VEGF) dependent key mechanism.

In contrast, excessive VEGF could be playing an etiologic role of VEGF-induced syndrome resembles peliosis hepatitis, suggested by the correlation between rising serum VEGF levels and the severity of the liver pathology, a rare human condition that is happened in the setting of advanced malignancies, high-dose androgen therapy and Bartonella henselae infection Wong et al. [26]. Similarly, liver peliosis occurred in follicular lymphoma with a rise in vascular endothelial growth factor and anaemia of inflammation De la Mano et al. [38]. Also, high levels of genetically induced or tumor-produced VEGF can alter the architecture of the adult liver, enhancing a liver ‘peliosis-like’ phenotype that was characterized by enlarged hepatic sinusoids, blood pooling, detached sinusoidal endothelial cells and a total disruption of normal liver architecture Belteki et al. 2005 and Wong et al. [26,39,40]. On conclusion, fas and loss of hepatocyte growth factor and vascular growth factor complex may be among the factor enhanced peliosis occurrence in liver.

References


