A Case Report of Hepatic Angiomyolipoma with Negative Premelanosome Marker Immunoprolife with Cirrhosis.

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Abstract: Angiomyolipoma (AML) is a relatively rare benign mesenchymal tumor that is frequently found in the kidney and, rarely, in the liver.[1] Hepatic AML is a rare, primarily benign mesenchymal tumor, composed of blood vessels, fat tissue, and smooth muscle cells [2]. Ishak reported the first hepatic AML in 1976 [3] and since then, there have been about 200 cases reported in the literature [8-13] and they have been increasing with improvement in imaging modalities, including ultrasonography (US), computed tomography (CT), magnetic resonance imaging (MRI), and fine-needle aspiration biopsy (FNAB)[4]. The hepatic AML may pose a diagnostic challenge clinically, radiologically, and pathologically because of its wide variation due to the different proportions of the three cell types which make up the tumor. In particular, in a region endemic for hepatocellular carcinoma, the diagnosis of AML by imaging modality can be difficult and frequently misdiagnosed as hepatocellular carcinoma. The definitive diagnostic study remains the histological examination coupled with immunohistochemical stains. A 44-year-old female, a known case of hemangioma liver presented with abdominal distension, loss of appetite and loss of weight. CECT revealed giant liver hemangioma. Enucleation of the tumour done. Histologically, tumour composed of dilated and thick walled arteries, adipose tissue and smooth muscles. Immunohistochemistry with HMB-45 done and found to be negative. Hence this case is presented for its rarity – THE MOST DIAGNOSTIC CRITERION [5] WHICH IS LACKING “HMB-45”

Keywords: hepatic angiomyolipoma, HMB-45

I. Introduction:
Hepatic angiomyolipoma (HAML) is a rare benign mesenchymal liver tumor first described by Ishak in 1976 [3]; it belongs to a group of perivascular epithelioid cell tumors called PEComa [7]. The tumor composed of blood vessels, smooth muscle, and adipose cells and due to the variety of predominance of these tissues, its patterns in imaging studies have resulted in a difficulty in diagnosis and misdiagnosis of the tumor as hepatocellular carcinoma (HCC) in some cases. [16,19]. Therefore, the preoperative correct diagnosis has been difficult; however, recent advances in imaging diagnosis through a combination of ultrasonography (US), computed tomography (CT), magnetic resonance imaging (MRI), and angiography and specific immunohistochemical analysis of this tumor using human melanoma black-45 antigen (HMB-45) staining have resulted in accurate diagnosis and it is reported that the current accurate preoperative diagnosis was made in 25%-52% of cases. [17,18] The majority of these tumors are believed to be clinically benign during a mean follow-up period of 6.8 years, [16] however, an increasing number of cases and aggressive changes including growth in size, recurrence after surgical resection, metastasis, and invasive growth pattern into the parenchyma and along the vessels have been reported.

II. Case History:
44-year-old female, a known case of hemangioma liver presented with complaints of abdominal distension, loss of appetite, loss of weight and early satiety. Per abdominal examination revealed massive hepatomegaly. Investigations revealed elevated Serum bilirubin, serum alkaline phosphatase and Prothrombin time and normal AFP levels. OGD – Normal. Contrast enhanced CT revealed 2 large hypodense with peripherally nodular, non-contiguous intense enhancement with central non-enhancing areas in right and left lobes of liver, largest measuring 22x15.6cms in right lobe – FEATURES SUGGESTIVE OF GIANT HEMANGIOMA OF LIVER WITH ASCITES.
Patient proceeded with Enucleation of the tumour. Intraoperatively, lesion is soft-firm in consistency involving segment 3,4a, exophytic in nature compressing stomach and transverse colon.
Grossly,

Received 2 soft tissue masses largest measuring 17x12.5x8cms and smallest measuring 12x11x8cms. External surface- grey brown and studded with tiny nodules. C/s: vaguely circumscribed solid brownish firm mass admixed with yellowish greasy areas and foci showing greenish glistening areas.

Histologically, Section shows liver parenchyma with adjoining areas show numerous thick walled vessels lined by endothelial cells, smooth muscle cells and adipose tissue. Adjacent liver tissue shows Cirrhosis.

Dilated blood vessels interspersed by smooth muscles
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This case is presented for its rarity.

III. Conclusion:

IV. Discussion:

Angiomyolipomas are rare in liver. They most often present at 30-40 years of age with female preponderance. Around 10% of tumors occur in the setting of tuberous sclerosis.[22,23] It is considered as a tumor of perivascular epitheloid cells and related tumors in other organs including renal angiomyolipoma, clear cell sugar tumor and lymphangioleiomyomatosis.[15] Patients present with features of abdominal pain, discomfort, malaise, fever, and anorexia. Rare occurrences have also been reported in many sites including the uterus, retroperitoneum, mediastinum, renal capsule, the hard palate, the nasopharyngeal cavity, the buccal mucosa, fallopian tube, vagina, penis, skin, abdominal wall, stomach, and spinal cord.[22,23]

Grossly, right lobe more often involved. They can grow as large as 36 cms in diameter. Cut surface - variably coloured depending on the content. Fat is yellowish colour and soft in consistency. Smooth muscle is tan white and firm. Hemorrhage may be present.[22,23] Histologically, composed of varying proportions of smooth muscle like cells, blood vessels and fat often in association with hematopoietic cells.[22,23]

According to the proportion of tissue components, Hepatic angiomyolipoma is classified into lipomatous (>70%), myomatous (<10%) and angiomatous.[8] The smooth muscle-like cells are prominent in liver lesions and consists of epitheloid or spindled cells that often surround or “spin off” vessels. The epitheloid cells are round to polygonal with abundant eosinophilic cytoplasm and large nuclei with prominent nucleoli. The cytoplasm may be oncocytic and condensed around the nucleus with a clear zone near the cell membrane giving “spider web” like appearance.

The vascular component is made up of thick walled arterial or venous-like channels admixed with thin walled venous like spaces.

The fatty component is made up of fat cells arranged singly, clusters or sheets.

Immunohistochemistry with HMB-45, smooth muscles are negative for the marker.

fat cells

Cirrhosis
Foam cells, Peliotic spaces lacking endothelial lining, dense lymphoid aggregates composed of T and B cells, Hemosiderin, melanin pigment and variable numbers of hematopoietic elements are also seen.[22,23]


HMB stands for Homatropine methyl bromide. Its target is unique pre melanosome related polypeptide. It stains cytoplasm of tumour cells in melanoma, PEComas, adrenocortical tumour and sex cord tumours of gonads.[21] HMB -45 is the diagnostic criterion for angiomyolipoma.[5] But in our case, since all the components of the tumour are evident morphologically, in spite of HMB-45 being negative, diagnosis of HMB-45 is made.

The lack of PMA expression in our patient’s case may be due aberrant antigenic expression. It is tempting to speculate that specific mutation(s) of the genes expressing premelanosome epitopes are missing in our patient’s case, although specific molecular genetic studies have not been performed for confirmation. PMA negative expression has been described even in melanoma cases, and electron microscopic studies need to be performed to confirm the presence of premelanosome structures.[14]

It is typically a benign lesion and resection is curative. Rare malignant variants with monotypic epithelioid variant, necrosis, marked pleomorphism and high mitotic activity.[16,20]

References:


[22] Goodblum. Surgical Pathology of gt, liver, biliary tree and pancreas. 2nd edition,pg 986