Abstract. Background/Aim: Epithelioid angiomylipoma of the liver is a rare benign mesenchymal tumor that usually presents in adult female patients. It most frequently occurs in the kidney, with the liver being the second most common site of involvement. Angiomylipoma belongs to a family of tumors arising from perivascular epithelioid cells, but in rare cases may also have cystic features. We report our experience via the first case of hepatic angiomylipoma treated by laparoscopic approach. Patients and Methods: We present the case of a 50-year-old female patient complaining of abdominal pain. Abdominal ultrasound (US) and Magnetic Resonance Imaging (MRI) revealed a 5 × 3 cm mass located in the left liver lobe. The tumor was resected with a laparoscopic approach. Microscopic examination of the tumor revealed hepatic angiomylipoma. Results: Twenty-seven months postoperatively, the patient remains fit and healthy. Conclusion: Angiomylipoma can be removed by laparoscopy.

Epithelioid angiomylipoma (AML) of the liver is a rare benign mesenchymal tumor that usually presents in adult female patients; nonetheless, several cases have also been reported among males. AMLs most frequently occur in the kidney, with the liver being the second most common site of involvement (1).

AMLs belong to a family of tumors arising from perivascular epithelioid cells (commonly called PEComas in the literature). Histological examination demonstrates that AMLs consist of adipose tissue, smooth muscle bundles, and thick-walled blood vessels in varying proportions, thus explaining the origin of their name.

Another characteristic feature of AMLs is the presence of epithelioid cells, particularly in the epithelioid variant of the tumor. Hepatic epithelioid AMLs may also have cystic features, but this is extremely rare (2).

In this report, we present the case of a 50-year-old female patient that was referred to our Department, after reporting discomfort in the right upper quadrant of the abdomen. Following a comprehensive initial work-up, a magnetic
resonance imaging (MRI) scan demonstrated a well demarcated hepatic mass that was surgically resected. Pathologic examination revealed the presence of a hepatic epithelioid AML with extensive cystic degeneration, a rare but, fortunately, benign finding. Subsequently, the patient was advised to return for periodic follow-up. At the time of the last follow-up visit (27 months postoperatively), the patient remained asymptomatic with no evidence of recurrence.

Case Presentation

We present the case of a 50-year old female patient that was referred to our department, after complaining of right upper quadrant abdominal pain for the past 4 months. The pain was not related to food intake and failed to respond to painkillers. The patient’s past medical history was unremarkable with no evidence of liver or biliary disease; furthermore, the patient reported no alcohol consumption. Clinical examination revealed tenderness in the right upper quadrant of the abdomen, as well as sensitivity to deep palpation in the right subcostal area. Laboratory examination demonstrated significant elevation of alanine aminotransferase (ALT) and aspartate aminotransferase (AST), as well as slightly decreased levels of total protein (Tp) and albumin (Alb). Alpha-fetoprotein (AFP), carcinoembryonic antigen (CEA) and carbohydrate antigen 19-9 (CA-19.9) levels were within normal limits. The patient underwent abdominal ultrasound (US) which revealed an isoechoic 5×3 cm mass in the left liver lobe. Subsequently, a contrast enhanced MRI was performed; it demonstrated a well-defined cystic mass in the left liver lobe with a hypointense T1 signal and a hyperintense T2 signal (Figure 1A and B). A heterogeneous enhancement was observed in the walls of the cystic mass; furthermore, the presence of adipose tissue was demonstrated in the mass’s cystic center (Figure 1C and D).

Figure 1. Angiomyolipoma located in the left liver lobe. A, B: On T1-weighted (A) imaging a sharply demarcated lesion with low signal intensity can be noted in the left liver lobe which shows a mildly hyperintense signal in T2-weighted imaging (B); C, D: The lesion is markedly hypervascular in the arterial phase (C) and remains hyperintense in the portalvenous phase (D).
The patient underwent exploratory laparoscopy, which revealed a solid tumor in the left hepatic lobe. With the aid of intra-operative US, the tumor’s borders and location with respect to the surrounding hepatic parenchyma and important vascular structures were carefully identified. No intrahepatic metastasis or satellite lesions were detected. Consequently, the decision was made to proceed with a left lateral hepatectomy (liver segments II and III) and concurrent cholecystectomy, both of which were performed laparoscopically. Following the successful completion of these procedures, a drainage tube was placed adjacent to the transected surface of the liver.

Gross examination revealed a circumscribed, smooth, multilobular tumor measuring $4 \times 4 \times 2.8$ cm. The cut surface of the tumor was spongiform and brownish in color, while the tumor wall demonstrated variable thickness. The tumor was resected intact and clear margins were achieved (R0 resection).

Microscopic examination of the tumor revealed multiple epithelioid cells with ample eosinophil cytoplasm, enlarged nucleus and prominent nucleoli. Interestingly, many of the epithelioid cells resembled ganglion cells; as noted above, the increased size of the nucleoli was their most remarkable feature. Neoplastic cells were arranged in broad solid sheets surrounding a prominent network of sinusoid-like vessels, filled with hemorrhagic fluid. Apart from epithelioid cells, the tumor also consisted of sinusoid-like vessels, spindle myoid cells and mature fat cells. Mitotic figures were not detected in any of these cellular populations, while focal areas of necrosis and hemorrhage were observed (Figure 2).

A resection margin width of 3 mm was achieved.

Immunohistochemical analysis revealed that epithelioid tumor cells were positive for HMB45, Melan-A (MART1) (Figure 3A and B), as well as for smooth muscle actin (SMA) (Figure 3C). On the other hand, the spindle myoid cells were positive for SMA. Both epithelioid and spindle cells were negative for CD10, HePar-1 (Figure 3D), S-100, epithelial membrane antigen (EMA), AFP, c-kit, chromogranin A and synaptophysin. The proliferation index was very low in all examined cellular populations.
The patient’s post-operative course was uneventful and all routine laboratory tests were within normal limits. The drainage tube was removed after the 4th post-operative day and the patient was discharged 8 days later. On follow-up examination, the patient remained asymptomatic and healthy to the present day. Computed tomography scans were also performed 6, 12, 18 and 24 months postoperatively; both imaging studies demonstrated no evidence of recurrence or other abnormal findings. A total of 27 months postoperatively, the patient remains fit and healthy.

**Discussion**

AMLs are mesenchymal tumors of benign nature, most commonly encountered in the kidney and closely associated with tuberous sclerosis (3). The liver is the second most common site of AML occurrence (1). Hepatic AML can present similarly to malignant tumors, such as hepatocellular carcinoma, both clinically and radiologically. Hepatic AMLs are difficult to diagnose radiologically, because of the variability of their growth patterns and their proximity to major hepatic vascular structures. However, the presence of mature adipose tissue may facilitate the accurate preoperative diagnosis of this rare clinical entity (4). It has been reported that hepatic AMLs are more commonly located in the right lobe (5). However, solitary involvement of the left lobe is also possible, as illustrated by the present case.

Hepatic AML was first described in 1976 by Ishak et al. (6) and is known to present with a wide variety of growth patterns. These variations reflect the distribution and degree of differentiation of myoid cells within the tumor and have been described by Tsui et al. (7). Several distinct AML histologic types have been identified, including the lipomatous, myomatous, angiomatous, trabecular, pelioid, inflammatory and mixed variants (8). Gross examination of hepatic AMLs usually reveals soft, solitary, well encapsulated tumors (4). The presence of cystic elements filled with hemorrhagic contents, as in the present case, is
very rare. Regardless of their site of occurrence, AMLs are generally composed of epithelioid and spindle myoid cells, mature adipose tissue and sinusoid-like vessels. The presence of multiple epithelioid cells within a tumor can be misleading and may render the differentiation of AML from hepatocellular carcinoma challenging (9).

In this report, we presented a rare case of hepatic epithelioid AML with extensive cystic degeneration in a 50-year-old woman. The tumor demonstrated the characteristic pathologic features of AML, such as the presence of spindle myoid cells, mature adipose tissue and sinusoid-like vessels; epithelioid cells were the predominant cell type. Immunohistochemistry was pathognomonic for AML, as epithelioid and spindle myoid cells were positive for melanocytic (HMB45) and myogenic (SMA) markers, but negative for epithelial markers (EMA).

The treatment of hepatic AML is mostly surgical. It is recommended that small lesions should be closely observed and treated conservatively, while larger lesions should be excised with clear margins. The present report is, to our knowledge, the first to describe the laparoscopic excision of a hepatic AML; a more detailed study of the efficacy of laparoscopic techniques in this setting should be considered.

Conclusion

AML is a rare, benign mesenchymal liver tumor that may present similarly to a hepatic malignancy. We herein presented a rare case of hepatic AML that was uncharacteristically located in the left liver lobe and demonstrated cystic degeneration with limited intracystic hemorrhage. AML should always be a part of the differential diagnosis in patients that present with solitary hepatic lesions; familiarity with the characteristic features of this uncommon clinical entity is essential to formulating an appropriate diagnostic and therapeutic strategy.

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