Management of perivascular epithelioid cell tumor of the liver: A case report and review of the literature

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Abstract. Perivascular epithelioid cell tumor (PEComa) is a rare tumor that originates from mesenchymal tissues. Cases of PEComa in the liver are extremely rare. The present study aimed to analyze the clinical features of PEComa in the liver and discuss its management. Here we report a 25-year-old female with multiple lesions of low density with poorly defined borders in the liver, as shown by a computed tomography (CT) scan. A partial hepatectomy was proceeded and PEComa was diagnosed by immunohistochemistry. No evidence of recurrence was observed during the one year follow-up. A total of 20 patients with hepatic PEComa, including one case from the present study and 19 cases that were reported in literature between June 2001 and December 2012, were reviewed and analyzed. The mean patient age was 43.4 years (range, 25-67 years) and the cases consisted of 18 female and two male patients. The tumor size ranged between 2.0x1.6 and 15.0x12.0 cm. Of the 20 patients, nine were asymptomatic and 11 had mild to significant complaints. Immunohistochemistry plays a key role in the diagnosis of PEComa. All the cases in this study were strongly positive for human melanoma black-45. A surgical resection is the gold standard for curative intent. All the patients underwent a surgical resection and none were administered perioperative chemotherapy or radiotherapy. In total, 13 of the 14 patients with follow-up information survived during the 8-36-month follow-up period and one patient succumbed due to recurrence two years after the surgery.

Introduction

Perivascular epithelioid cells (PECs) were first proposed in 1992 by Bonetti et al (1) and classified by the World Health Organization in 2002 (2,3). The PEC tumor (PEComa) family is composed of epithelioid angiomyolipoma (AML), clear-cell ‘sugar’ tumors, lymphangioleiomyomatosis, clear-cell myomelanocytic tumors of the falciform ligament/ligamentum teres and rare clear-cell tumors of other anatomical sites (4-6). PEComa is mainly composed of eosinophilic and clear epithelioid cells, which are commonly arranged as small nests that are associated with variably-sized vessels (5-7). The predominant site of origin for PEComa is the uterus, but the tumor may also be found in the falciform ligaments, prostate and kidney. However, cases in the liver are extremely rare (2,8). Hepatic PEComa has a marked female predominance and possesses no specific symptoms (9). The diagnosis of PEComa is based on its pathological characteristics, including epithelioid cells without adipocytes or abnormal blood vessels, and on immunohistochemical evidence, including melanocytic and smooth muscle markers (8). Surgery is the only effective method to result in a long survival time (6,9).

In the present study, one case of hepatic PEComa is described. Furthermore, 19 cases from the literature, in which 11 patients were diagnosed with hepatic PEComa and eight with hepatic epithelioid angiomyolipoma, are reviewed.

Case report

Presentation and laboratory examinations. A 25-year-old female who presented with an abdominal mass, which was revealed by ultrasonography, was admitted to the Second Affiliated Hospital, Zhejiang University School of Medicine (Zhejiang, Hangzhou, China) in December 2011. The past history and physical examination were normal. The laboratory examinations revealed a slightly elevated level of carbohydrate antigen 19-9 (CA199; 38.8 kU/l; reference range, <37 kU/l), the levels of alanine aminotransferase (ALT; 7 U/l), aspartate aminotransferase (AST; 14 U/l), serum creatinine (SCR; 49 µmol/l), blood urea nitrogen (BUN; 3.80 mmol/l), α-fetoprotein (AFP; 2.2 µg/l) and carciinoembryonic antigen (CEA; 0.9 µg/l) were within the reference ranges. Hepatitis B and hepatitis C virus screenings were negative.

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Diagnostic techniques. A plain computed tomography (CT) scan demonstrated multiple lesions of low density with poorly-defined borders in hepatic segments III (0.5x0.5 cm), IV (5.4x5.5 cm) and VII (1.8x1.5 cm). The delayed phase showed mild intensity lesions. However, the intensity of the lesions increased significantly in the contrast-enhanced phase (Fig. 1A). A liver magnetic resonance imaging (MRI) scan revealed that the lesions had a medium signal intensity on the T1-weighted image and a slightly high signal on the T2-weighted image. The enhancement of intensity of the lesions was also observed in the contrast-enhanced phase of MRI (Fig. 1B).

Treatment. During the laparotomy, three well-encapsulated tumors located in segment III, IV and VII were identified. No portal or inferior vena cava vein invasion or distant metastasis was observed. A partial hepatectomy of the liver neoplasms was performed using the Pringle maneuver.

Pathology and immunohistochemical analysis. The masses in segments III and IV were pathologically identified as hemangioma, while the mass in segment VII was revealed to be composed of polygonal morphology cells, which are similar to epithelial cells. However, the latter mass did not contain lipocytes or abnormal blood vessels. Immunohistochemistry further revealed that the mass in segment VII was strongly positive for human melanoma black-45 (HMB-45; +++; Fig. 2A), smooth muscle actin (SMA; +++; Fig. 2B) and vimentin (++). The mass was positive for CD34, but negative for S-100, creatine kinase (CK), epithelial membrane antigen (EMA), desmin, AFP and Ki67. A diagnosis of PEComa of the liver was confirmed based on the immunohistochemical analysis. The patient recovered well and was discharged one week after surgery. No evidence of recurrence was observed during one year of follow-up.

Study approval. Approval for this study was obtained from the ethics committee of Zhejiang University and informed consent was provided by the patient.

Literature review

Identification of patients with hepatic PEComa. The Chinese BioMedical Literature Database, the China Hospital Knowledge Database and the Wanfangbase were searched between January 2001 and December 2012. Repeated studies were carefully screened and rejected from further analysis to avoid over-representation. The final diagnosis of each case was confirmed using pathological and immunohistochemical results. The data on the clinical features, tumor characteristics, pre-operative imaging and outcome of treatment of 19 cases were collected and analyzed along with the data of the present case.

Presentation. The mean patient age was 43.4 years (range, 25-67 years) in the 20 patients. PEComa was shown to have a marked female predominance (18 females and two males; Table I). The chief presenting complaints were abdominal pain (5/20), abdominal discomfort (5/20) and abdominal distension (1/20). Of the 20 patients, nine were asymptomatic and discovered the mass incidentally during a physical examination. The majority of the tumors were localized in the right lobe (16/20) of the liver. The tumor size ranged between 2.0x1.6 cm and 15.0x12.0 cm. Of the 20 tumors that were analyzed, seven were >5 cm in size at the time of presentation.

Pathological findings and immunohistochemistry. Histological examination revealed that the tumor was highly
cellular, with large round or polygonal cells with abundant cytoplasm and clear cell boundaries. Immunohistochemistry revealed that the tumor cells in all cases were positive for HMB-45 (20/20) and melan-A (9/9). The cells were positive for SMA (14/16), vimentin (12/13) and S-100 (7/13) in the majority of the cases (Table II).

Treatment. All 20 patients underwent surgery following admission. Three patients were treated with a hemihepatectomy, three with a segmentectomy and 10 with a partial hepatectomy of the liver neoplasms. However, no patients were administered chemotherapy or radiotherapy.

Follow-up. The follow-up data was available for 14 patients and the follow-up time ranged between 8 and 36 months. One patient succumbed due to recurrence at two years post-surgery and the others survived without recurrence or metastasis.

Discussion

The pathogenesis of PEComa, although discussed in previous studies, remains controversial. Kenerson et al (10) demonstrated that tuberous sclerosis complex 1/2 (TSC1/2) inactivation and mTOR hyperactivation were present in non-TSC AMLs and extrarenal PEComas using immunohistochemistry and western blot analysis. In particular, mTOR hyperactivation may be studied in such lesions using immunohistochemical detection of p70S6K, which is a marker of mTOR activity (6).

Bing et al (11) also showed that epithelioid AMLs harbored p53 mutations in certain cases.

Hepatic PEComa occurs most commonly in females (7), and symptoms of hepatic PEComa usually show no specificity. In the present study, 11 out of 20 patients with hepatic PEComa had mild to significant non-specific complaints. The remaining nine patients were asymptomatic. The majority of the patients (16/20) had solitary lesions in the right lobe, which was consistent with a previous study by Parfitt et al (7).

Ultrasonography, CT and MRI are most frequently employed for the pre-operative diagnosis of PEComa. Previous studies have considered hypervascularity and

| First author, year (ref.) | Gender/age, years | Symptom | Size, cm | Location, lobe |
|---------------------------|-------------------|---------|----------|----------------|
| Present case              | F/25              | None    | 2.5x1.0  | Left           |
| Qiu et al, 2008 (19)      | F/67              | Abdominal pain | 15.0x12.0 | Right         |
| Deng et al, 2011 (20)     | M/66              | None    | 3.0x3.5  | Left           |
| He et al, 2011 (21)       | F/35              | None    | 3.5x3.0  | Right          |
| Han et al, 2008 (22)      | M/44              | None    | 2.0x1.6  | Right          |
| Li et al, 2007 (23)       | F/56              | Abdominal distension | 5.0x4.0 | Left         |
| Chen and Li, 2009 (24)    | F/37              | Abdominal pain | 5.0x4.0  | Right         |
| Zhang and Wang, 2012 (25) | F/55              | None    | 3.0x3.0  | Right         |
| Chen, 2009 (26)           | F/36              | Abdominal pain | 7.0x5.0  | Right         |
| Chen, 2009 (26)           | F/45              | Abdominal discomfort | 5.5x4.0 | Right         |
| Liu et al, 2010 (27)      | F/32              | None    | 5.5x5.5  | Right         |
| Zou et al, 2011 (28)      | F/54              | Abdominal discomfort | 6.0x5.0 | Right         |
| Xu et al, 2001 (29)       | F/35              | None    | 2.0x2.0  | Right         |
| Lin et al, 2002 (30)      | F/30              | Abdominal discomfort | 3.6x3.1 | Right         |
| Liu et al, 2008 (31)      | F/31              | Abdominal pain | 8.0x6.0  | Right         |
| Zhu et al, 2010 (32)      | F/26              | None    | 5.0x3.0  | Right         |
| Gao et al, 2012 (33)      | F/59              | Abdominal pain | 6.0x5.0  | Right         |
| Wang et al, 2007 (34)     | F/46              | None    | 4.0x4.0  | Right         |
| Shi et al, 2010 (35)      | F/41              | Abdominal discomfort | 5.5x4.0 | Left         |
| Shi et al, 2010 (35)      | F/48              | Abdominal discomfort | 8.0x6.0 | Right         |

M, male; F, female.
arteriovenous connections to be a feature of PEComa in contrast-enhanced CT (8,12,13). An MRI scan revealed that the PEComas were significantly and heterogeneously enhanced in the arterial phase, but less enhanced in the portal venous and delayed phases (14,15), which may also effectively rule out a diagnosis of hepatocellular carcinoma with a fibrotic capsule (15). Another commonly used diagnostic method is contrast-enhanced ultrasonography, which possesses the features of an early influx of the contrast agent into the tumor and a rapid drainage of arterial blood to the veins (15). However, the accuracy of pre-operative diagnosis is low, partly as a result of the variable imaging appearances due to the varying proportion of the components, including the smooth muscle cells, adipose tissue and vessels, and the rarity of the tumor. In the present case, three lesions of the liver shared similar imaging features, but only one lesion was confirmed to be a PEComa. In the cases that were reviewed in this study, three patients were pre-operatively diagnosed with focal nodular hyperplasia (including the present case), four patients were diagnosed with hepatic carcinoma and 12 patients could not be differentiated.

The final diagnosis of PEComa depends on the pathology and immunohistochemistry. Martignoni et al (6) defined PEComa as a tumor that is composed of solely of cells with an epithelioid appearance, which are closely associated with dilated vascular channels and contain clear eosinophilic cytoplasm, but do not contain lipocytes or disordered blood vessels. Almost all the PEComas were identified to be strongly positive for melanocytic markers (HMB-45 and/or melan-A) and smooth muscle markers (SMA and/or desmin) (2,5,16). In the present study, the tumor cells of the 14 patients were positive for HMB-45 and SMA, but in the case reported by Zou et al (28), only HMB-45 and melan-A were positive and SMA was not stained.

PEComa has been reported to exhibit benign behavior in the majority of the literature. Since the first case of malignant hepatic PEComa presented by Dalle et al (17), several malignant cases have been reported (7,17,18). In 2005, Folpe et al (16) reviewed 26 cases of soft tissue and gynecological origins and raised seven criteria to evaluate the malignancy of PEComa: i) A tumor size of >5 cm; ii) infiltration into the surrounding normal tissue; iii) a high nuclear grade; iv) hypercellularity; v) high mitotic activity (>150 high-power field); vi) coagulative necrosis of the tumor; and vii) vascular invasion. A malignant PEComa is considered to have two or more of the features that are listed. Tumors with nuclear pleomorphisms, multinucleated giant cells only or those of >5 cm in size are considered as neoplasms of uncertain malignant potential. In the present study, one case of malignant hepatic PEComa was also reviewed in which the patient was identified to have a recurrence two years after the surgery.

Surgical resection with an adequate margin remains the gold standard for the treatment of hepatic PEComa (6,9,14), particularly in malignant cases. In the present study, all patients underwent surgical treatment. The malignant patient underwent a hemihepatectomy due to the enormous tumor size. Hemihepatectomies were also performed in another two patients, who were misdiagnosed with hepatocellular carcinoma prior to the surgery. Generally, chemotherapy and radiotherapy does not indicate an improved survival time (6). However, several studies have shown promising treatment results, including a study of rapamycin, which is an inhibitor for mTOR. Rapamycin had a positive effect on renal angiomylolipoma (5). If rapamycin is able to yield the same effect on other PEComas, this would provide the rationale for lesions that are composed of PEs. Of the 14 patients in the present study, 13 were alive during the 8-36-month follow-up period and one succumbed due to recurrence. However, Parfitt et al (7) noted that PEComa may also demonstrate recurrence following a long period of time (nine years). Thus, the prognosis of PEComa remains unpredictable, and it is necessary to perform long-term follow-up studies for every case.

In conclusion, the diagnosis of hepatic PEComa depends on the pathological observations. Surgical resection of the tumor appears to be necessary for a cure, particularly for malignant tumors. Although adjuvant chemotherapy or radiotherapy are not significant in the treatment, several studies that have used drugs to treat PEComa have shown promising results. In the present study, the tumors were mostly benign and the prognosis following surgical resection was good.

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