Case Series

Primary Hepatic Angiosarcoma: A Rare Liver Malignancy – Varying Manifestations but Grave Prognosis

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Keywords
Primary hepatic angiosarcoma · Hepatic angiosarcoma · Intra-tumoral hemorrhage

Abstract
Primary hepatic angiosarcoma (PHA) is a rare mesenchymal liver tumor, accounting for 0.1–2% of primary liver malignancies. The clinical presentations of PHA are variable, from asymptomatic to liver failure or complicated with tumor rupture. The diagnosis of PHA is difficult due to the lack of specific clinical manifestation and investigation results, which can be confused with other liver tumors resulting in late diagnosis. However, there is currently a paucity of effective therapeutic approaches. We advocate early diagnosis with radiological imaging and histopathology because most of them are diagnosed in late-stage and carry a grave prognosis. Surgical resection remains the mainstay of treatment, which can significantly prolong survival. Chemotherapy, including transarterial chemoembolization, is an option for palliative treatment. Unfortunately, molecular treatment has limited efficacy and liver transplantation is also not recommended due to high rate of recurrence. We present a case series of four patients with biopsy-proven PHA which had distinct presentations and clinical courses.
Introduction

Primary hepatic angiosarcoma (PHA) is a rare hepatic tumor, originating from endothelial and fibroblastic tissue, primarily made up of vessels and composed of abundant vasculature. It represents only 0.1–2% of all primary liver malignancies [1]. PHA commonly occurs in ages from 60 to 70 years. The clinical presentations are variable, from asymptomatic to liver failure; most patients have nonspecific symptoms at initial presentation that usually mimic other chronic liver diseases and are difficult to distinguish from other liver malignancies, resulting in late diagnosis and delayed treatment. These symptoms include abdominal pain, fatigue, weight loss, jaundice, and anorexia. Physical examination can reveal hepatomegaly, ascites, or jaundice [2]. Few patients present with hemoperitoneum secondary to tumor rupture, which was reported in 17–27% of cases [3, 4]. The differential diagnosis of PHA are hepatocellular carcinoma, cholangiocarcinoma, metastatic carcinoma of liver, and hepatic hemangioma. Furthermore, there is a lack of specific laboratory findings for diagnosis of PHA, include liver function test or tumor markers. Imaging study and pathological features remain the key to aid diagnosis. On computed tomography (CT), the dominant tumors usually show heterogenous enhancement, suggesting central necrosis or hemorrhage and delayed progressive enhancement [5–7]. Pathologically, PHA is composed of pleomorphic atypical mesenchymal cells with elongated nuclei, and the tumor exhibits its own disorganized anastomotic vascular channels. The areas of necrosis and hemorrhage are commonly reported in histological findings [3], when more specific immunohistochemical markers for PHA including ERG expression, CD34, CD31, and factor VIII can lead to definite diagnosis [8]. Unfortunately, PHA is associated with grave prognosis. Surgical resection remains a curative treatment; however, it is difficult to perform because 80% of patients have advanced stage at diagnosis. Chemotherapy and transcatheter arterial chemoembolization (TACE) are considered a palliative treatment. Liver transplant is not recommended due to a high rate of recurrence and rapid progression of tumor with survival after liver transplant less than 7 months. The majority of patients have medial survival less than 1 year, even after receiving treatment. The common causes of death are liver failure and hemoperitoneum from spontaneous tumor rupture [4, 9, 10].

There has been a paucity of data and significant challenges regarding the diagnosis, treatment, and prognosis of this tumor. Many physicians remain unaccustomed to the clinical features of PHA. We present the cases of four patients with biopsy-proven PHA, which had distinct presentations and clinical courses.

Case Presentation

Case 1

An 81-year-old man was admitted due to massive upper gastrointestinal hemorrhage for 2 h. He had epigastric pain and early satiety for 3 weeks. His past medical history was unremarkable except for heavy alcohol drinking, 1–2 bottles a day for over 30 years. He denied a history of exposure to thorotrast or vinyl chloride. Physical examination revealed mildly pale, icteric sclerae, hepatomegaly with a liver span of 14 cm, parotid gland enlargement and palmar erythema. Other examinations were unremarkable. Laboratory investigations on admission showed white blood cell (WBC) count 6,210/dL, hemoglobin (Hb) 5.8 g/dL, platelet count
283 × 10³/dL, aspartate transaminase (AST) 116 U/L, alanine aminotransferase (ALT) 73 U/L, alkaline phosphatase (ALP) 192 U/L, albumin 2.7 g%, total bilirubin (TB) 5.74 mg%, direct bilirubin (DB) 5.3 mg%, prothrombin time (PT) 16.2 s (control 12.4), and international normalized ratio (INR) 1.23; serum alpha-fetoprotein (AFP) was within normal range.

Esophagastroduodenoscopy was performed and revealed a 3-cm large ulcer with elevated border and necrotic tissue in the antrum of stomach; no varix was seen. The tissue biopsy was performed to exclude malignant ulcer. Liver ultrasonography revealed a cirrhotic liver and diffused multiple echogenic nodules size 7–9 mm in both lobes with a large echogenic mass 3.3 cm in size with a halo. A further CT scan demonstrated multiple large infiltrative masses with few arterial enhancements, no venous washout, size 1–4 cm in both lobes, osteolytic foci at the thoracic spine level T10 and T11, and small subpleural nodule at the right lower lung. Liver biopsy was performed for definite diagnosis. The specimen consisted of a few pieces of gray tan irregular soft tissue and disclosed the angiosarcoma; immunohistochemical study showed positive staining for CD34 and factor VIII. The specimen of gastric ulcer biopsy also revealed metastasis of angiosarcoma.

He was treated with a paclitaxel dose of 175 mg/m². Unfortunately, 5 days after receiving chemotherapy, he developed febrile neutropenia with septic shock. His relatives refused further treatment and invasive procedures, and he died in the following days.

Case 2
A 55-year-old woman with a medical history of type 2 diabetes mellitus, presented with right upper abdominal pain and significant weight loss for 5 months, and progressive jaundice for 2 months. She denied a history of carcinogen exposure. Physical examination showed hepatomegaly 4 cm below the right costal margin with a liver span of 15 cm. Liver examination described a rubbery to firm consistency, blunt edge, and smooth surface. Other findings were unremarkable except mild icteric sclerae. The complete blood count revealed: WBC 9,820/dL, Hb 12.0 g/dL, and platelet count 98 × 10³/dL. The liver function test showed: AST 44 U/L, ALT 20 U/L, ALP 449 U/L, TB 3.81 mg%, DB 2.57 mg%. The serum AFP was 2.16 ng/mL (normal <7 mg/mL).

Multiphase CT scan demonstrated diffused multiple arterial enhancing infiltrating nodules varying in size, scattered throughout the entire liver, with ascites and mesenteric fat reticulation suspicious of carcinomatosis peritonei. Laparoscopic liver biopsy was performed for diagnosis, and the specimen consisted of a piece of light brown irregular soft tissue. Histopathological findings showed the presence of diffused proliferation of abnormal small vessels with diffused hemorrhage. The abnormal vessels showed enlarged endothelial cells with nuclear atypia. Immunohistochemical study showed positive staining for CD34 and factor VIII in the endothelial cells. Histopathological study was consistent with PHA (Fig. 1).

She received a total of five cycles of paclitaxel with a 20% dose reduction. The following posttreatment CT scans showed progression of the disease with peritoneal metastasis, and she later decided to stop treatment. She survived for 19 months and 14 months after symptom onset and treatment initiation, respectively.
**Case 3**

A 47-year-old woman without past medical history was admitted due to acute right upper abdominal pain for 8 h and syncope prior to admission. She had a history of taking oral contraceptive pills 4–6 months per year, for 7 years. Physical examination revealed vital signs: BP 100/60 mm Hg, pulse rate 115/min. Her body mass index was 33.6 kg/m². Abdomen examination showed marked tenderness at the right upper quadrant; the liver was palpated 8 cm below the costal margin with a liver span of 20 cm with mild tenderness, no splenomegaly, and no shifting dullness.

Laboratory investigations showed: WBC count 8,200/dL, Hb 7.4 g/dL, and platelet count 592 × 10³/dL, AST 34 U/L, ALT 23 U/L, ALP 335 U/L, TB 1.22 mg%, DB 0.73 mg%; coagulogram was in normal limit, AFP was 2.72 ng/mL.

The emergency CT scan revealed acute rupture of liver tumor by evidence of acute clot and acute hematoma. She was provisionally diagnosed with ruptured hepatic adenoma and underwent emergency hepatic embolization. The angiography finding revealed a large hepatic mass at the anterior aspect of the right hepatic lobe, receiving arterial supply from the anterior branch of the right branch of the hepatic artery. Eventually, embolization was performed to stop bleeding. The scheduled CT scan 2 weeks after embolization showed a larger diameter of hemorrhagic mass from 20.7 to 23.0 cm, lysed perihepatic hematoma, without distance metastasis (Fig. 2).

Four weeks after embolization, she underwent extended right heptectomy with wedge diaphragm and primary repair. The operation demonstrated a huge liver mass with intratumoral hemorrhage, displaced IVC, and mediastinum. The specimen consisted of a liver tumor size 28 × 20 × 14 cm and weighing 2,300 g. The tumor appeared to be a light-brownish rubbery mass with a massive hemorrhage. The histopathological finding was compatible with angiosarcoma (French system grade III) and free tumor margin. The immunochemical staining was positive for CD34 and factor VIII, consistent with PHA (Fig. 3).

The follow-up CT scan was performed after surgery, and it showed multiple new nodules about 0.5–2 cm in the left lobe liver, a heterogenous enhancing mass size 3.7 cm in the right subdiaphragmatic space; peritoneal thickening was also noted. Thus, the physicians planned to treat her with chemotherapy, doxorubicin. However, she refused chemotherapy treatment because of her poor clinical condition, and she passed away 4 months later.

**Case 4**

A 69-year-old previously heathy woman presented with left upper quadrant pain for 2 months. She described her intermittent colicky abdominal pain, each attack lasting 10 min. She also noticed a weight loss of 3 kg in a week. Physical examination showed palpated liver 4 cm below the right costal margin and 6 cm below the xiphoid; the surface of the liver appeared to be lobulated with firm consistency; no hepatic bruit was heard.

The laboratory investigation showed: WBC count 5,080/dL, Hb 10.5 g/dL, and platelet count 592 × 10³/dL, AST 34 U/L, ALT 23 U/L, ALP 335 U/L, TB 1.22 mg%, DB 0.73 mg%; coagulogram was in normal limit, AFP was 2.72 ng/mL.

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For a definite diagnosis, she underwent real-time sonographic guidance percutaneous tumor biopsy. The specimen consisted of light brown soft tissue. The histopathological finding revealed a malignant vascular tumor, immunohistochemistry staining positive for CD 34 and factor VIII. These findings are consistent with angiosarcoma.

The patient had planned to have chemotherapy. Unfortunately, 2 weeks after diagnosis, she was admitted to the hospital due to severe abdominal pain. Investigations revealed a ruptured tumor, evidence of disseminated intravascular coagulopathy, and liver failure. Thus, she denied invasive therapy and passed away 1 week later.

The clinical manifestation, tumor feature, and treatment outcome of the four patients are summarized in Table 1.

Discussion

PHA is a liver tumor originating from endothelial and fibroblastic tissue, accounting for 0.1–2% of all primary liver malignancies [1]. Despite its rarity, PHA remains the most common malignant mesenchymal cell tumor of the liver [2]. PHA generally develops in the sixth to seventh decade of life and is predominantly found in males with a male-to-female ratio of 2–4:1 [11]. According to an epidemiological study from the United States, 25% of PHA were associated with known etiologies. Thorotrast application in angiography was linked to PHA in the past decade with the average latency period from exposure to onset of about 20 years. Vinyl chloride, in plastic fabrication, has also been shown to have an established association with PHA with the hazard ratio approximately 10- to 15-fold. Besides, other agents including arsenic androgen, anabolic steroids, cyclophosphamide, phenelzine, and copper are proposed as possible carcinogens [2, 9]. However, these cases are rare and lack explicit associations.

The presentations of PHA are variable, ranging from asymptomatic to acute liver failure. Most of the patients have nonspecific symptoms that mimic chronic liver diseases, with late diagnosis as the consequence. These symptoms include abdominal pain, fatigue, weight loss, and anorexia [2]. The reported physical examinations of PHA comprised hepatosplenomegaly, jaundice, ascites, and rarely hepatic bruit due to its vascular nature. Few patients, approximately 9%, present with symptoms secondary to metastasis. Hemoperitoneum is a fetal complication secondary to tumor rupture, which was reported in 17–27% [3, 4].

Moreover, rare presentations of PHA in cases of acute liver failure, high-output heart failure, esophageal varices, hemothorax, and disseminated intravascular coagulopathy from Kasabach-Merritt syndrome have recently been reported [12].

For investigation, almost all patients (97%) undergo a nonspecific abnormal liver function test. Accretion of ALP is the most common abnormal finding. Hyperbilirubinemia may occur in cases of advanced disease. Cytopenia is common in PHA, with 54% of patients displaying thrombocytopenia. While anemia may reflect spontaneous tumor rupture or microangiopathic hemolytic anemia, there is no specific tumor marker corresponding to PHA [4].

The lack of specific clinical presentations and laboratory findings emphasizes the diagnostic value of radiological imaging. Conventional ultrasound has reported a limited yield in diagnostic evaluation. The mass typically demonstrated different echogenicity depending on necrosis or hemorrhage nature.
On imaging, similar findings appear on dynamic CT and magnetic resonance imaging (MRI). The dominant tumor usually showed heterogeneous enhancement suggesting central necrosis, hemorrhage, and fibrotic change. Delayed progressive enhancement but incomplete venous and delayed phase of enhancing lesions is observed without a centripetal filling pattern. In comparison to hemangioma, the enhancing pattern of PHA is a peripheral rim with central septal-like or linear progression, while a hemangioma-enhancing pattern is centripetal nodular enhancement [5–7].

On T1-weighted MRI, the irregular high signal intensity of the dominant mass suggests the presence of hemorrhage. On T2-weighted images, there is increased signal intensity and a septum-like or rounded area of low intensity. The hyperintense signal indicates the area of hemorrhage or necrosis, while the hypointense signal represents hemosiderin deposition and fibrous solid portion. On dynamic contrast-enhanced MRI, the enhancement pattern shows the progressive enhancement of lesion as a dynamic CT scan [5–7].

Lastly, relevant angiographic findings comprise numerous mass lesions with fluffy staining and early pooling of contrast media that progressively increase over time. The most prevalent feature of the angiogram is intense peripheral stain late in the arterial phase.

The diagnosis of PHA remains problematic owing to lack of obvious clinical and investigation features, which are typically analogous with other vascular tumors. A definite diagnosis requires histopathology via liver biopsy. Many reports recommend open liver biopsy because of better visualization and easier hemostasis, whereas percutaneous biopsy, a simpler method, is a safer and faster procedure without significant complications [13]. Post-biopsy hemorrhage is the primary concern in PHA because of its vascularity. Bleeding occurrence was reported in 9%, which can be managed by blood transfusion and hepatic arterial angiembolization.

Pathologically, PHA consists of four characteristics; multiple nodules, large dominant mass, mixed mass with nodule, and diffusely infiltrating micronodular tumor. The tumor has a pale white-yellow-gray color, a spongy appearance, and an ill-defined margin [3].

PHA is composed of pleomorphic atypical mesenchymal cells, polyhedral, and sometimes spindle-shaped [8]. Neoplastic cells contain prominent chromatin, atypical hyperchromic, and elongated nuclei with frequent mitoses. This tumor exhibits its own disorganized anastomosis vascular channels, ranging from dilated sinusoid or cavernous spaces to slit-like freely anastomosing vessels, formed by spindle-shaped cells [3]. Areas of necrosis, hemorrhage, and calcification are reported in 80% of specimens.

There are no specific immunochemistry markers. A recent report stated that the most sensitive and specific marker is ERG expression, with a 100% sensitivity, followed by CD34 (87.5%), CD31 (87.2%), and factor VIII (41.7%), respectively [8].

For the treatment of PHA, there is a paucity of effective therapeutic approaches. Surgical resection remains a promising treatment for PHA. Radiosurgery with R0 resection is the sole curative treatment. Adjuvant chemotherapy provides further survival benefit with reported median survival up to 17 months [10]. Liver transplantation is contraindicated in PHA owing to high recurrence and poor prognosis regardless of liver transplantation.

Chemotherapy is considered a palliative treatment in patients who are not surgical candidates. Many chemotherapy regimens have been demonstrated in various reports, none of which appear to be superior. Salvage chemotherapy with 5-fluorouracil/carboplatin/doxorubicin/ifosfamide improves survival in 50% of patients, based on its antiangiogenic properties.
The limited efficacy of molecular treatment including bevacizumab, sorafenib, and sunitinib cannot be recommended. TACE is effective in patients with a dominant mass. Reports have shown that TACE with either lipiodol/cisplatin or lipiodol/mitomycin/Adriamycin is beneficial in 50% of patients [14, 15]. Radiotherapy is not a treatment option for PHA due to the intrinsic radio-resistant property of tumor cells.

PHA carries a grave prognosis. Postdiagnosis median survival without treatment is approximately 1–6 months, with 3% surviving longer than 2 years [4, 9, 10]. For patients who underwent surgical resection either with or without adjuvant chemotherapy, the median survival is approximately 17 months [10]. Curative surgery is difficult to perform in more than 80% of cases because of the advanced stage at diagnosis [4]. The most common cause of death is liver failure, accounting for 50% of cases, followed by hemoperitoneum from spontaneous tumor rupture (25%). Other causes of death are metastasis disease, infection, renal failure, and heart failure (3%) [4, 10]. Distant metastasis makes up more than 60% of cases, which are diagnosed post-mortem. The most common site of metastasis is the lungs, spleen, and bone, respectively. Complications are reported for 17–27% of individuals [4].

In conclusion, PHA is a rare, hepatic, and malignant tumor, which physicians may not recognize as it mimics other liver tumors resulting in late diagnosis. Clinical presentations are variable from mild to severe symptoms, and fetal complications may be present. Spontaneous tumor rupture or liver failure may commonly occur. Diagnosis of the disease in the early stage with CT or MRI and histopathology provides survival benefit. However, prognosis is still poor despite chemotherapy treatment.

Acknowledgements

The authors acknowledge Dr. Poowadon Wetwittayakhlang for providing histological pictures for these cases.

Statement of Ethics

Informed consent was obtained for this case series report. The names of the subjects are not mentioned in this case series, and no identifying image has been included for these cases.

Disclosure Statement

The authors have no conflicts of interest to declare.

Funding Sources

There are no funding sources to report.
Author Contributions

N. Rujeerapaiboon wrote the manuscript, and P. Wetwittayakhlang wrote and supervised the writing of the manuscript and provided the clinical information of all cases. Both authors read and approved the final manuscript.

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Fig. 1. Histopathological examination of tumor tissue in case 2. **a** Histopathological study (hematoxylin-eosin stain, ×40) showed the tumor cells (T) grow along sinusoids adjacent to hepatic cords (L). **b** The tumor cells appear abundant, poorly defined cell borders, pleomorphic with hyperchromatic nuclei. The abnormal vessels also showed enlarged endothelial cells with nuclear atypia (×400). The Immunohistochemical study showed positive staining for CD 34 (**c**) and factor VIII (**d**).
Fig. 2. Multiphase CT scan of liver in case 3 demonstrates a large hemorrhagic hepatic mass 15 × 20 × 23 cm in size with an enhanced solid part at the anterosuperior part on plain phase.
Fig. 3. Histopathological finding of primary hepatic angiosarcoma in case 3. a Histological examination of tumor tissue (hematoxylin-eosin stain, ×40) showed an area of the tumor tissue (T) infiltrated in the liver tissue (L). b The tumor cells present diffuse proliferation of abnormal small vessels and enlargement of endothelial cells with nuclear atypia (×400). The immunohistochemical study of the same tissue showed positive staining for CD 34 (c) and factor VIII marker (d), which indicated hepatic angiosarcoma.
Fig. 4. Multiphase CT scan of the liver in case 4 illustrates multiple scattered hypervascular infiltrative masses and nodules, about 1–9 cm in size in an almost entire enlarged liver. The masses have progressive enhancement on portal venous phase, and most of them have surrounding ground-glass opacity which is a typical finding for hemorrhage in angiosarcoma.
Table 1. Summary of clinical features, tumor characteristics, treatments, and survival outcomes in the four cases

| Case | Age, years | Sex | Clinical manifestations | Tumor characteristics | Extrahepatic metastasis | Tumor rupture | Treatment | Survival time from symptom onset |
|------|------------|-----|-------------------------|-----------------------|------------------------|--------------|-----------|---------------------------------|
| 1    | 81         | M   | massive upper GI bleeding, abdominal pain, jaundice | multiple infiltrative mass | spine, pleura, stomach | no          | paclitaxel 20% dose reduction | 1.5 months |
| 2    | 55         | F   | abdominal pain, jaundice, weight loss | multiple scattered nodules | peritoneal | no | paclitaxel 20% dose reduction | 19 months |
| 3    | 47         | F   | intraperitoneal hemorrhage | single dominant mass with hemorrhage | none | yes | surgical resection | 6 months |
| 4    | 69         | F   | abdominal pain, weight loss, hepatomegaly | diffuse multiple mass with hemorrhage | spine, lungs | yes | best supportive care | 2 months |