Imaging findings of splenic hamartoma

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AIM: To assess CT and MR manifestations and their diagnostic value in splenic hamartoma with review of literatures.

METHODS: We described a woman who was accidentally found to have a splenic tumor by ultrasound of the abdomen. CT and MR findings of this splenic hamartoma were proved by pathology retrospectively.

RESULTS: The CT and MR findings in this case included a ball-like mass with homogeneous mild-hypodensity lesions on non-enhanced CT scans or isointensity on T₁-weighted images and mild hypointensity on T₂-weighted images, progressive homogeneous enhancement on multiple-phase spiral CT and MR enhanced scans, and isodense enhancement on delayed post-contrast CT scans and obvious hyperintensity relative to the spleen on delayed MR images.

CONCLUSION: Splenic hamartoma has some specific radiological features. However, the diagnosis of this disease must be based on clinical features and confirmed by pathology.

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INTRODUCTION

Splenic hamartoma is a rare benign vascular tumor[1]. Up to now, less than 50 cases in Chinese literature or about 160 cases in the world have been reported after the first description by Rokitansky in 1861.

Imaging features of splenic hamartoma have been described by several researchers at computed tomography (CT), magnetic resonance imaging (MRI) and sonography[2-8] and the imaging appearance of these lesions is considered as nonspecific and a histopathological confirmation is often required[9-11].

This paper described the CT and MRI features of splenic hamartoma in a 40-year-old patient with review of the literature.

CASE REPORT

A 40-year-old woman was accidentally diagnosed having a splenic tumor by ultrasound of the abdomen. She did not complained about fever, fatigue, abdominal pain, or weight loss was complained. She denied any history of hepatitis or tuberculosis. Physical examination was entirely normal. There was no evidence of jaundice, peripheral lymphadenopathy or hepatosplenomegaly. Her hemoglobin was 125 g/L, the number of white blood cells was 7.3×10⁹/L and the number of platelets was 176×10⁹/L. Both stool and urine routine tests and hepatic and renal function tests were normal. Chest x-ray was also normal.

Spiral CT of the abdomen before contrast medium administration revealed a 3.8-cm diameter, homogeneous mild-hypodensity lesion within the spleen (Figure 1A). During hepatic artery phase following bolus injection of intravenous contrast, the lesion showed a slightly homogeneous enhancement (Figure 1B). During the portal venous phase and hepatic parenchymal phase, the lesion showed a progressively homogeneous enhancement. On delayed images up to 5 min after-contrast injection, the tumor was isodense with the spleen (Figure 1C). CT diagnosis was a splenic hemangioma.

MRI of the abdomen showed a 3.6 cmx3.8 cmx4.2 cm ball-like mass with isointensity on T₁-weighted images (Figure 2A) and mild hypointensity on T₂-weighted images (Figure 2B). The lesion demonstrated a diffuse heterogeneous enhancement on images obtained early after contrast medium administration (Figure 2C) and became more uniformly enhanced on the portal venous phase and hepatic parenchymal phase. It was hyperintense compared to the spleen on the delayed images (Figure 3). The MR examination yielded a diagnosis of splenic hemangioma. A 4.0-cm diameter tumor was found in the spleen during operation. The pathologic diagnosis was a splenic hamartoma (Figure 4).

Figure 1  CT scans of splenic hamartoma. A: Homogeneous mild hypodensity lesion within the spleen found by unenhanced CT scan, B: Mild-homogeneous enhancement of the mass found by enhanced CT scan on hepatic artery phase, C: Isodense tumor with normal spleen on delayed enhanced CT images.
DUSCUSSION

Hamartoma of the spleen is a rare benign lesion and the diagnosis is difficult to make preoperatively. Histologically it is composed of an aberrant mixture of the normal tissue components of the spleen, so hamartoma of the spleen is often called splenoma, splenadenoma or nodular hyperplasia[2,3]. Splenic hamartoma occurs most commonly in adults. About 14.3% of the reported cases of splenic hamartoma occurred in pediatric patients[12]. Most patients were asymptomatic, they were incidentally found during imaging studies, laparotomy or autopsy[2,3]. Our case was accidentally discovered by ultrasound examination of the abdomen. Symptomatic splenic hamartoma was rare but nearly half of splenic hamartoma pediatric patients had symptoms[12]. A minority of these lesions had hematologic symptoms such as pancytopenia, anemia, and thrombocytopenia[12-10]. Spontaneously ruptured splenic hamartoma has been reported[17,18]. Symptomatic splenic hamartoma with renal, cutaneous abnormalities or portal hypertension and heterotopic ovarian splenoma were described accidentally[19,21].

A few radiological findings in splenic hamartoma have been described[2-8]. Sonography was a more sensitive modality than CT in demonstrating the lesion, which showed hyperechoic masses with cystic components occasionally[2]. But hypoechogenic splenic mass was found and color Doppler sonography showed blood-flow signals inside the mass in a recent report[4]. CT could reveal splenomegaly and homogeneous or heterogeneous low-density or isodense masses with calcification[2,5,6] or fatty components, which are characteristic CT findings. Dense spreading enhancement on dynamic CT and prolonged enhancement on delayed post-contrast scans were noted in singular mass[15]. But low-density masses relative to the spleen were seen in multiple splenic hamartomas after contrast medium administration[2]. The CT findings in this case were similar to those in Ohtomo’s report[5], which included a homogeneous mild-hypodensity lesion on non-enhanced scans, a progressive homogeneous enhancement on multiple-phase spiral CT enhanced scans and an isodense enhancement on delayed post-contrast scans.

There were two types of MRI findings in splenic hamartomas, fibrous and non-fibrous splenic hamartomas. Histopathologically, fibrous splenic hamartomas had a dominant fibrous tissue and MRI showed isointensity or hyperintensity on T1-weighted images, hypointensity on T2-weighted images[3]. We consider hypointensity on T2-weighted images is one of the common MRI findings in splenic hamartomas. Non-fibrous splenic hamartomas are more common and MRI showed isointensity on T1-weighted images, hyperintensity on T2-weighted images[6,8]. Both of the tumors demonstrated diffuse heterogeneous enhancement on the hepatic artery phase or early dynamic contrast-enhanced scans, which became more uniformly enhanced on delayed images[8,9]. This case belonged to fibrous type and the MRI findings were similar to those in Fernandez-Canton’s report besides obvious hyperintensity relative to the spleen on delayed images[3]. We agree with that diffuse progressive enhancement and prolonged enhancement were the characteristic radiological findings[3].

In short, the following clinical features and radiographic findings may suggest the diagnosis of splenic hamartomas: (1) asymptomatic and incidental findings in adults; (2) possible association with hematologic symptoms such as pancytopenia, anemia, and thrombocytopenia or spontaneous rupture of splenic mass; (3) splenic mass with calcification or fatty components on plain CT, and isointensity on T1-weighted images, heterogeneous hyperintensity or hypointensity on T2-weighted images; (4) dense spreading enhancement and obviously prolonged enhancement on postcontrast CT and MRI. Though splenic hamartomas have some clinical and CT features, the final and exact diagnosis depends on histopathologic examination.
Splenectomy is the only curative treatment for splenic haemangiomata. They have similar clinical and radiological findings. Both splenic haemangiomata and splenic hamartomas are seen on plain CT. In addition, splenic lymphoma and metastases are also considered to be different from splenic hamartomas. Most of splenic lymphomas and metastases are multiple, secondary lesions and often have extra-splenic lymphoma. Splenic lymphomas usually have a history of primary extra-splenic malignant neoplasms and hepatic metastases. Both splenic lymphomas and metastases seldom have dense spreading enhancement or prolonged enhancement on post-contrast CT and MRI.

REFERENCES


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