Abstract. Upper gastrointestinal bleeding caused by a gastric hemangioma is a relatively rare occurrence. We report the case of a 65-year-old woman who suffered a sudden onset of hematemesis. Endoscopy revealed a 4x3 cm mass located in the gastric fundus. Abdominal contrast-enhanced CT revealed the shadow of enhancing linear blood vessels located in the gastric fundus. Based on her clinical appearance and the laboratory results, the patient was diagnosed with gastric hemangioma. In the laparotomy, a proximal gastrectomy was performed. The final diagnosis of cavernous hemangioma arising from the gastric fundus was confirmed by postoperative pathological examination.

Introduction

Hemangioma appear in various parts of the human body, including the liver, spleen, colon, retroperitoneum, adrenal glands, soft tissues, extremities, central nervous system and mediastinum (1). Gastric hemangioma, a rare tumor occurring mostly in the antrum, was first described by Lammers in 1893 (2). It represents approximately 1.7% of all gastric benign tumors and 20% of unknown hemorrhage. In this case report, we present the clinical presentation, diagnosis and treatment of a cavernous hemangioma that arises from the gastric fundus.

Case report

The patient is a 65-year-old Chinese female who suffered a sudden onset of hematemesis with approximately 800 ml blood loss in a total of 10 h. When she was admitted to hospital, she was in a state of pre-shock with the complaint of muscle weakness and dizziness. She had no past history of diabetes, hypertension, hepatitis or bronchial asthma. The patient did not take any medication, including over-the-counter non-steroidal anti-inflammatory drugs (NSAIDs), and was not taking herbal supplements. On abdominal examination she felt discomfort only in the upper abdomen, without tenderness, and she was hemodynamically unstable. Routine blood tests, including hepatic and renal function tests, revealed a hemoglobin level of 56 g/l, a platelet count of 93×10^9/l, elevated blood urea nitrogen (BUN) of 10.25 mmol/l, total protein of 53.9 g/l, albumin of 30.8 g/l, serum creatinine of 60.9 µmol/l, lactate dehydrogenase (LDH) of 199 U/l, uric acid of 121.0 µmol/l, and an erythrocyte sedimentation rate (ESR) of 31 mm/h. Based on the patient's clinical appearance and the laboratory analysis, she was diagnosed with acute upper gastrointestinal bleeding and hemorrhagic shock.

An urgent endoscopy following active resuscitation revealed a 4x3 cm mass located in the gastric fundus with active hemorrhage at the surface (Fig. 1A). The bleeding was controlled with an injection of 1 ml 1:10,000 adrenaline (Fig. 1B). The patient underwent prompt treatment with vigorous intravenous (IV) rehydration, blood transfusion for worsening anemia, and IV ranitidine therapy. Following stabilization, abdominal computed tomography with its periphery enhanced by the contrast material in the delayed phase revealed the shadow of enhancing linear blood vessels located in the gastric fundus (Fig. 1). No abnormalities were revealed in the laboratory data of the tumor markers, which are listed as: CA19-9, 6.39 KU/l (<35.00); CA242, 3.41 KU/l (<20.00); CA125, 2.27 KU/l (<35.00); CA15-3, 2.28 KU/l (<35.00); NSE, <1.0 ng/ml (<13.00); CEA, 3.25 ng/ml (<5.00); ferritin, 20.10 ng/ml (male<322.00, female<219.00); β-HCG, <0.03 MIU/ml (<3.00); AFP, 5.68 ng/ml (<20.00); HGH, 2.00 ng/ml (<7.50). In the laparotomy, a soft mass located in the gastric fundus was found to vibrate when pressure of the hand was applied. A proximal gastrectomy was performed. The postoperative period was uneventful and the patient was discharged 10 days after the surgery. The final diagnosis of cavernous hemangioma arising from the gastric fundus was confirmed by postoperative pathological examination.

Discussion

Kaijser has classified gastrointestinal hemangioma pathologically as multiple phlebectasia, cavernous hemangioma, capillary hemangioma and angiomatosis (3).
Abdominal cavernous hemangiomas usually originate in the liver, but occasionally present in the stomach with the same origin and mechanism as other cavernous angiomas, which have been considered as a congenital, benign and abnormal development. Findings of previous studies have shown that cavernous hemangiomas are congenital hamartomatous lesions that originate from the mesodermal remnant tissues. These hemangiomas are composed of large dilated blood vessels and contain large blood-filled spaces that are caused by dilation and thickening of the walls of the capillary loops (4). Due to the thin walls of blood vessels, gastric cavernous hemangiomas (GCH) are prone to rupture with rapid blood flow. In this case report, we highlight the rarity of this lesion and the difficulties in diagnosing it preoperatively. Most gastrointestinal tract hemangiomas are of the cavernous type and upper gastrointestinal bleeding is the most common symptom (5). Gastric hemorrhages are common clinical emergencies, which often directly involve the surgeon in diagnosis and treatment; among these, rare cavernous hemangiomas deserve particular attention.

Endoscopy may serve to establish the diagnosis in those cases where, due to the hemangioma's small size or unfavorable location, radiological examination fails to detect them (2). Usually, emergency endoscopy is a means of diagnosis and treatment, but this was not effective in the present case since the delicate tissue tended to bleed. Furthermore, it is possible that this characteristic did not allow a correct tissue sample to be collected for pathological examination, which yielded...
a negative result. Radiological examination suggests the possible diagnosis, and the best definitive diagnostic procedure is CT scanning and MRI, which demonstrate the location and relationship of the lesion to neighboring structures as the preoperative reference of resectability. The lesion appears either as enhancing linear blood vessels or caputmedusae, a radial orientation of small vessels that resemble the hair of Medusa from Greek mythology. However, when the blood vessel signal is weak, CT scanning and MRI are incapable of distinguishing GCH from mesenchymal tumors. However, GCH may be misdiagnosed as a stromal tumor. Surgical treatment is a definitive modality, and recurrence following complete resection has not been reported thus far. The final diagnosis in the present case was obtained by definitive histopathology.

In conclusion, we report a rare case of cavernous hemangioma originating from the gastric fundus. It may cause diagnostic difficulties preoperatively as biopsy is not an option due to the submucosal location of the tumor and hemorrhagic factor. However, in the present case, we were able to obtain the result of enhancing linear blood vessels by contrast-enhanced CT scanning prior to the surgery, which assisted us in making a primary diagnosis. Therefore, abdominal images should be examined on a regular basis. Although this disease is benign with a lower recurrence following total resection, we nevertheless suggest the requirement for long-term follow-up to assess treatment outcome.

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References