An uncommon mass of the esophagus: esophageal hemangioma

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Case report

A 73-year-old woman complaining of retrosternal pain and swallow discomfort over the past 1 month was admitted to our hospital. She denied any dysphonia, regurgitation or weight loss. Physical examination was unremarkable. The hematological and biochemical profiles were normal. Barium meal studies revealed a smooth oval-shape mass with broad pedicle in the posterior wall of the upper esophagus (Figure 1). The patient refused to endoscopic investigation because of irregular heart rhythms. Further chest CT scan revealed a soft tissue mass with punctuate calcification and narrowing of the lumen to around 40% of the original size. The lesion showed moderately in homogenous enhancement after contrast injection (Figure 2). There was no lymphatic node adenopathy in the mediastinum. Based on her symptoms and related radiological findings, the patient was firstly suspected with esophageal hemangioma and underwent thoracoscopic enucleation. Intraoperative photography showed deep-red lesion with smooth surface (Figure 3) and the mass was completely resected under the thoracoscopy. Histopathological findings were consistent with esophageal haemangioma (Figure 4). Post-operative course was uneventful, and the patient’s symptom of retrosternal chest pain was completely resolved after surgery. She remained asymptomatic at 2-year follow-up.

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from esophageal vascular tumor can be massive or even fatal. Barium esophagogram findings are nonspecific, only showing either a well-defined lobulated intramural mass, pedunculated intraluminal mass or an infiltrating annular mass. Endoscopic ultrasound is the most sensitive and specific method for assessing the location, depth and size of an esophageal benign tumor [1]. On CT examination, an esophageal hemangioma usually appears as a well-defined soft tissue mass within the esophageal wall. Phleboliths associated with inhomogeneous enhancement after contrast injection are considered to be pathognomonic findings for this uncommon tumor as described in our case. Although esophageal hemangiomas are rare, they can usually be identified by above specific imaging features aided with clinical symptoms and endoscopic findings, instead of biopsy.

As for treatment, enucleation of hemangiomas is considered as treatment of choice. However, option includes esophagectomy or tumor enucleation by thoracotomy or an endoscopic approach [2]. Recently successful treatment has been reported with endoscopic sclerotherapy [3]. In the case of rich vasculature tumors, endoscopic resection can be troublesome due to uncontrollable bleeding. On the basis of broad-based and rich vascular mass in our patient, thoracoscopic removal of esophageal hemangioma was successfully performed with excellent outcome.

References