A case of leiomyoma of the oesophagus complicated by superior vena cava obstruction and associated eosinophilia

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Summary
Superior vena cava obstruction occurring as a complication of leiomyoma of the oesophagus has not been reported before. Such a case is recorded which was associated with striking eosinophilia, a feature previously noted in cases of uterine leiomyomas.

Case report
A 35-year-old merchant seaman had complained of intermittent epigastric pain, water brash and heartburn for approximately a year. These symptoms were usually associated with drinking bouts in which he frequently and freely indulged, but which he had recently curtailed. He had 3 episodes of haematemesis during the year, each being associated with drinking. For 6 months he had noticed increasing dysphagia for solids and he occasionally regurgitated solid food. In March 1978 he had a barium meal examination which showed a smooth compression of the oesophagus measuring 7 cm x 5 cm posteriorly and to the right side of the aortic arch (Fig. 1). This investigation was repeated with tomography and confirmed a large extrinsic impression on the right side of the oesophagus. No hilar gland enlargement was identified. At this time his white cell count was 8.2 x 10^9/l, with a normal differential. He was awaiting further investigation when in May he presented as an emergency with a 6-day history of increasing swelling of his left arm, left hand and his face.

On examination, there was pitting oedema of the dorsum of his left hand and a firm swelling of the whole of his left arm. Both supra-clavicular fossae and the suprasternal notch were obliterated. There was swelling over the neck, face and anterior chest wall, and moderate conjunctival suffusion. The jugular veins were distended to the level of the ear lobes and were non-pulsatile. The retinal veins were

Fig. 1. Barium meal examination—the leiomyoma is seen compressing the right side of the oesophagus.
engorged but there was no papilloedema. General examination was normal and in particular there was no lymphadenopathy. His peripheral pulses were normal, blood pressure 120/70 mmHg in both arms, and there were no abnormal signs in the chest or abdomen.

Investigations at that time were as follows: Hb, 16.6 g/dl; white cell count, 23.1 x 10⁹/l; (15% eosinophils, 66% neutrophils, 15% lymphocytes, 1% monocytes and 3% myelocytes).

Bone marrow cytology showed a highly cellular marrow with a striking degree of eosinophilia, constituting 65% of the total cells of which eosinophil myelocytes constituted 20%, eosinophil metamyelocytes 40% and mature eosinophils 40% of the total eosinophil count. His Mantoux test was negative at a dilution of 1/10000.

Management and progress

It was decided to treat his superior vena caval obstruction with steroids. He was given prednisone 40 mg/day, and the signs in his face, neck and left arm resolved completely over the next 48 hr. The eosinophilia increased over this period with the WCC rising to 36.5 x 10⁹/l, of which 28% were eosinophils. Upper gastro-intestinal endoscopy was performed after the superior vena caval obstruction had settled. This revealed a smooth swelling approximately 3 cm in diameter, impinging on the oesophageal lumen at 25 cm. The mucosa overlying it was normal and there was no pulsation visible. At 35 cm there was a sliding hiatus hernia. The endoscope passed easily into the stomach which was normal in appearance.

A right thoracotomy was performed (K.B.). There was no mediastinal lymphadenopathy but there was a round, shiny tumour, 6.5 cm in diameter, within the right antero-lateral wall of the oesophagus. The trachea was deviated to the right and the tumour was bulging forwards closely adjacent to the junction of the innominate veins, theazygos vein and the superior vena cava (Fig. 2). The tumour was easily enucleated from the oesophageal muscle leaving the mucosa intact. Treatment with prednisone was stopped after 2 weeks.

Histology of tumour

Macroscopic appearance – a rounded encapsulated tumour, rubbery in consistency. The cut surface was glistening and white with areas of degeneration at both poles.

Microscopic appearance – the appearance was that of a leiomyoma with areas of necrosis. There were many focal and perivascular collections of eosinophils (Fig. 3).

The patient made an uneventful recovery. Post-operative barium swallow showed normal oesophageal motility with no evidence of leakage. The patient was soon swallowing normally. His differential blood count returned to normal. A superior vena cavaogram with injection of contrast into both arms was performed 2 weeks after surgery and this demonstrated patent central veins with no evidence of invasion, displacement or thrombus. He remains

Fig. 2. Diagram showing position of leiomyoma as seen at right thoracotomy.
perfectly fit, with no recurrence of his superior vena caval obstruction and his peripheral blood count is entirely normal, 6 months after removal of the oesophageal leiomyoma.

**Discussion**

Leiomyomas are the commonest benign tumours of the oesophagus (Seremetis et al., 1975; Mansour, Hatcher and Huan, 1977). Symptoms, if any, are usually confined to the local effects of pressure on the oesophageal lumen, such as dysphagia and chest pain. Acid reflux, usually seen in cases with associated hiatus hernia, as in this case, is also a common complaint. Weight loss, presumably due to dysphagia, is a less common symptom. Complications of oesophageal leiomyomas are rare, but malignant change, haemorrhage and asphyxias have all been reported (Seremetis et al., 1975). So far as the authors know, this is the first case in which potentially lethal obstruction of the superior vena cava has been reported. No other lesion could be found to account for the obstruction either radiologically, endoscopically or at surgery.

It was initially suspected that the patient had a mediastinal malignancy, possibly a lymphoma, in view of the peripheral blood eosinophilia. The prompt response of the obstruction to steroids strengthened these suspicions, but whether this response was coincidental or due to a direct local effect on the tumour and/or associated oedema of the surrounding tissues, is not known. No other cause for the eosinophilia was discovered despite an exhaustive search. He was not taking any drugs known to cause eosinophilia.

The most common haematological abnormality associated with leiomyomas, iron-deficiency anaemia excepted, is polycythaemia (Payne, Woods and Wrigley, 1969). A leucocytosis in the presence of an infected and/or necrotic leiomyoma sometimes occurs (Buka, 1965). Although well recognized in association with malignancy, eosinophilia is only rarely associated with benign tumours (Isaacson and Rapaport, 1946; Murray, 1953). Buka (1965) reported 2 cases of uterine leiomyomas associated with eosinophilia which resolved after hysterectomy. In neither case was the eosinophilia as marked as in the present case. The extensive eosinophilic infiltration of the tumour itself is also interesting in this case with respect to the peripheral blood and bone marrow eosinophilia. In the cases reported by Buka,
no areas of acute or chronic inflammation were present in the uteri, nor was there any collection of eosinophils in the leiomyomas themselves, and no report is made of areas of necrosis, in contradistinction to the histological findings in the present case. The stimulus for the eosinophilia in all 3 cases must remain conjectural but, like the polycythemia sometimes associated with leiomyomas, the eosinophilia has promptly resolved when the leiomyomas were removed (Rothman and Rennard, 1963). Eosinophilic infiltration of leiomyomas is not a feature which has hitherto been noted but eosinophilic infiltration of necrotic malignant tumours has been reported (Isaacson and Rapaport, 1946). It is thought unlikely, however, that in the case of leiomyomas, the necrosis alone is responsible for the eosinophilia, as necrotic degeneration, often associated with haemorrhage, is a common event in leiomyomas of the uterus and eosinophilia is not usually seen in these cases. Buka speculates that in his cases the eosinophilia is due to an auto-immune response to the leiomyomas which is manifested by the eosinophilia. It is not clear how this auto-immune response may be mediated but it is an attractive hypothesis to explain the puzzling association between leiomyomas and eosinophilia. The present authors can only conclude, as Buka does, that leiomyomas should be included in the differential diagnosis of any unexplained eosinophilia.

It appears, therefore, that the present case is one of leiomyoma of the oesophagus complicated by superior vena cava obstruction and associated with massive eosinophilia involving the tumour itself, the bone marrow and the peripheral blood.

References
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