Gastrointestinal Stromal Tumor of the Esophagus: Report of a Case

Özofagusun Gastrointestinal Stromal Tümörü: Bir Olgu Sunumu

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Abstract
Gastrointestinal stromal tumors are rare neoplasms to be thought to arise from mesenchymal cells of the gastrointestinal tract. Gastrointestinal stromal tumors (GIST) of the esophagus are well documented but are very much rarer than gastrointestinal stromal tumors of the stomach and small bowel. We describe a case of GIST of the esophagus that was resected with wide surgical resection.

Keywords
Gastrointestinal Stromal Tumor; Esophagus; Surgery

Özet
Gastrointestinal stromal tümörler (GIST) gastrointestinal sistem mezenkimal hücrelerinden gelişen nadir tümörlerdir. Özofagus kaynaklı GIST ise mide ve ince bağırsakla karşılaştırıldığında oldukça nadirdir. Bu olgu sunumunda geniş cerrahi reseksiyonla çıkarıldığımız bir GIST olgusunu sunuyoruz.

Anahtar Kelimeler
Gastrointestinal Stromal Tümör; Özofagus; Cerrahi
Introduction
The term gastrointestinal stromal tumor (GIST) refers to all mesenchymal tumors in the gastrointestinal tract [1]. These tumors demonstrate a pathobiology and clinical behavior different from those of smooth muscle and Schwann cell tumors. GISTs account for 0.1% to 3% of all tumors in the gastrointestinal tract. The majority of GISTs occur in the subdiaphragmatic gastrointestinal tract, but small number of cases has been in the esophagus [2, 3]. Here in we present a patient with a esophageal GIST treated with wide surgical excision.

Case Report
51 years old male patient had dysphagia and stomach ache for 6 months. We performed endoscopic examination and found external compression at 4 cm cranial to the gastroesophageal junction (Figure-1a). The computerized tomography showed a mass surrounding the esophagus at the same level (Figure-1b). We performed a right thoracotomy. We found a mass that began 4 cm below the azygous vein and lied distally. The mass was settled between muscular and mucosal layers of esophagus. We opened the muscular layer and completely resected the mass that was 3x7 cm in diameter and had thin capsule and a smooth border (Figure-1c,d). Result of frozen section was benign esophageal tumour so that after enucleating the muscular layers were sutured one by one over the mucosal layer. We discharged the patient at postoperative 3rd day. No mitosis was detected during pathological examination at 50 times magnification. Ki67 labeling index was 2/1000. CD117, Vimentin, CD34 (Figure-2), Desmin and SMA were positive and S100 was negative. The mass had the diagnosis of middle degree malign esophageal gastrointestinal stromal tumor. The follow up at 3rd month was uneventful.

Discussion
Gastrointestinal stromal tumors (GISTs) are uncommon mesenchymal tumors that arise in the wall of the gastrointestinal tract (GI). They account for approximately 0.1% to 3% of GI neoplasms [4]. GISTs were previously thought to be smooth muscle neoplasms, and most were classified as leiomyoma or leiomyosarcoma [2].

In 1983, Mazur and Clark challenged the longstanding concept that most mesenchymal tumors of the stomach were of smooth muscle origin, and introduced the concept of stromal tumor [3-5]. With the advent of immunohistochemistry and electron microscopy, it became apparent that GISTs might have myogenic features, neural attributes or characteristics of both muscle and nerve. Two thirds of these tumors arise from the stomach, 25% arise from the small intestine and 5% arise from the esophagus [2]. GISTs occur in an older patient population (50-60 years) [2, 3]. Approximately one half of patients with an esophageal GIST are asymptomatic, the remainder exhibit symptoms, which may include dysphagia, retrosternal pain, pyrosis, cough, odynophagia and weight loss [1]. Our case was 51 year – old male and had dysphagia and stomach ache for 6 months. Most esophageal GISTs are endocentric (intraluminal polypoid mass); on endoscopy the mucosa is usually intact and the mass appears as a rounded, smooth, raised lesion rarely showing central umblication or ulceration [1]. Esophageal GISTs originate from between the walls of the esophagus. They are kinds of proliferation of spindle cells or epithelioid cells [5]. The majority of GISTs are benign (60-80%). The most consistent prognostic factors are site of presentation and tumor size [3]. Kimiyashi suggested a criterion for differentiation between benign and malignant GISTs; hemorrhage or necrosis, the diameter of the tumor > 5cm, Ki-67 labeling index > 3%. If the tumor has only one of the items above it is malignant. If none of the items above can be found, then it is benign. The diameter of our mass 3x7cm, Ki-67 labeling index was 2/1000 and no mitosis was detected during pathological examination at 50 times magnification, so that our mass had the diagnosis of middle degree malign esophageal gastrointestinal stromal tumor.

GIST; there was focal positivity for pancytokeratin marker and for CD56 and diffuse strong positivity for CD117 and CD34. CD117 is sensitive and specific . Although CD34 is a sensitive immunochemistry marker of GISTs, CD34 is expressed in 60-70% cases of GISTs [5]. Our mass had positivity for CD34, CD117.

These tumors seem to be resistant to chemoradiation. Because of the lack of any effective alternative therapies, surgical resection should be considered for all patients with GISTs [2].

References