A Rare Cause of Hemoptysis in Childhood: Tracheal Capillary Hemangioma

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INTRODUCTION
Primary tumors of the trachea are extremely rare, with an estimated incidence of approximately 2.7 new cases per million per year, and usually malignant in adults [1]. Lobular capillary hemangioma (LCH) is a benign tumor, more commonly seen in children, with a distinctive lobular arrangement of capillaries in edematous and fibroblastic stroma [2]. The usual sites of tumor are the lips, nose, oral cavity, and tongue. It typically presents with nonspecific clinical symptoms, such as cough and hemoptysis. Stridor is observed with subglottic localization of the tumor. Radiologic studies and bronchoscopy are usually sufficient for diagnosis. Intervventional bronchoscopic techniques can treat these lesions and avoid aggressive surgical approaches. Mills et al.[3] reviewed 639 cases of vascular lesions of the oral cavity and upper respiratory tract. They found only 73 cases with LCH and no case with localization in or below the larynx. There is very limited literature on tracheal LCH [1]. We present here the diagnosis and management of a rare case of tracheal LCH that was successfully treated with interventional bronchoscopy.

CASE PRESENTATION
A 12-year-old boy was admitted to our clinic due to recurrent hemoptysis. His symptoms started 2 years ago with approximately 100 mL of hemorrhage in a day after coughing. He also mentioned that a small amount of hemoptysis recurred whenever he had a respiratory tract infection. There was nothing in his history to indicate causes of hemoptysis, such as tuberculosis, recurrent lower respiratory tract infections, intravenous or oral drug use, or antibiotic therapy. He had no pets. Physical examination showed no abnormalities. His hematologic parameters were as follows: activated partial thromboplastin time: 24.7 sc; prothrombin time:15.2 s-58.2%, INR: 1.22; within normal limits). Other chemical laboratory test results were normal, as were chest radiography findings (Figure 1). Sputum acid-fast bacilli and Löwenstein-Jensen cultures were negative. Thorax computed tomography showed a polypoid lesion on the left lateral wall of the proximal trachea (Figure 2). Rigid bronchoscopy revealed a reddish polypoid lesion with a smooth surface on the left lateral wall localized in the proximal one-third of the trachea. The lesion was attached to the tracheal wall with a short pedicle (Figure 3a). The lesion was excised with electrocautery snare and was taken using biopsy forceps. The tracheal wall was coagulated with argon plasma coagulation (APC) (Figure 3b). Histologic examination revealed a subepithelial lobular arrangement of proliferating capillary vessels, dilatation, and congestion; thus, the diagnosis of LCH was confirmed (Figure 4a, b). Upon follow-up, the patient had no hemoptysis episodes. On bronchoscopic examination at the 6-month follow-up, no relapse or other abnormality was observed except for a small scar on the previous biopsy site (Figure 3c). Written informed consent was obtained from the parents of the patient for this case presentation.
DISCUSSION

The most frequent causes of hemoptysis are infectious diseases, malignant tumors, cardiovascular diseases, and other inflammatory disorders in adults. LCH has seldom been reported in the English literature [4]. Chest X-ray findings are generally normal. LCH is frequently diagnosed while researching etiology of cough and hemoptysis with thorax CT and bronchoscopy [5]. Bronchoscopic appearance is also nonspecific and mimics adenoid cystic carcinoma, tracheal adenoma, and carcinoid tumors [6]. definitive diagnosis is made with bronchoscopic biopsy. Our patient was an adolescent whose chest X-ray findings were normal, but he had tracheal lesion on thorax CT, that was taken to determine the etiology of hemoptysis.

Pathogenesis of capillary hemangiomas (CH) is unknown. Bartonella infections are implicated in CHs of the skin. Minor trauma, such as endotracheal intubation and tracheostomy, has been reported to play a role in the etiology, but it is not clear because of the small number of cases in the literature [7]. There is no trauma history in our case. CHs can be successfully treated with various methods such as cryotherapy, yttrium aluminium garnet (YAG) laser, topical or intralesional steroids or neoplastic agents (vincristine), oral propranolol, and surgical excision. However, these treatments are mostly used for cutaneous, oral, and subglottic hemangiomas [8,9]. Although case reports are limited in number, successful results with different bronchoscopic techniques were obtained in the treatment of tracheal CHs.

Rameau et al.[10] successfully treated children presenting with symptomatic large tracheal hemangioma using K-potassium titanyl phosphate laser ablation, and they suggested this method for tracheal lesions. Excision of tumor with bronchoscopic forceps has also been reported as a successful method [2,4,5,7]. In our case, after the tracheal lesion was excised with electrocautery snare, excised tumor was taken with forceps and tracheal wall of the lesion was cauter-
ized using APC. After six months, control bronchoscopy revealed minimal scarring with no sign of recurrence.

In conclusion, we would like to remind that tracheal CH is an extremely rare cause of hemoptysis and radiological findings may be normal. So, it should be kept in mind for differential diagnosis. These exceptionally rare tumors also can successfully treated with interventional bronchoscopic methods.

**Informed Consent:** Written informed consent was obtained from the parents of the patients who participated in this case.

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