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Solitary myofibroma of the oropharynx causing airway obstruction in an adult

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Abstract
Myofibromas are benign neoplasms believed to be the most common fibrous proliferation of childhood. We present an unusual case of a 44-year-old woman who developed acute airway obstruction from a myofibroma in the oropharynx and accordingly required emergent tracheotomy tube placement. Serial laser excisions to adequately remove the entire lesion while maintaining pharyngeal structure and function were performed, and the patient was successfully decannulated. To date she has remained free of signs and symptoms of recurrence. Although rare in adults, solitary myofibromas should be considered in the differential diagnosis of any subcutaneous or submucosal head and neck lesion. Moreover, clinicians treating adult and pediatric patients with known solitary or multicentric forms of myofibroma should be aware of its potential for airway obstruction. Patients found to have a pharyngeal myofibroma should be managed with airway stabilization, surgical excision with preservation of speech and swallow function, and close postoperative monitoring for recurrence.

Introduction
Myofibromas are histologically benign neoplasms most frequently seen in the pediatric population in both solitary and multicentric forms. They may present in different locations, although they are often found in the oral cavity in both children and adults. These lesions are typically painless and slowly growing, and they follow an innocuous course. Myofibromas of the head and neck have not been previously described in the literature as presenting with acute airway obstruction and necessitating an emergent surgical intervention. To our knowledge, this is the first report of a solitary myofibroma of the oropharynx in an adult causing airway distress and requiring a tracheostomy.

Case report
A 44-year-old woman presented to our institution with a 3-month history of worsening dysphagia and position-dependent dyspnea. Fiberoptic examination revealed an oropharyngeal mass that completely obstructed visualization of the distal airway. Magnetic resonance imaging (MRI) demonstrated a 3.0 x 1.7 x 1.4-cm pedunculated lesion of the right base of the tongue and epiglottis, with pronounced enhancement with contrast and hyperintensity on fat-suppression images (figure 1). The patient was urgently taken to the operating room for biopsy, followed by subtotal resection to optimize the airway and preserve function. Histopathology revealed the lesion to be a myofibroma (figure 2). After airway stabilization, the patient was scheduled for elective resection in 2 weeks and discharged home. Having failed to return for her scheduled surgery, the patient presented again 3 months later in respiratory distress. She required tracheotomy tube placement to stabilize her airway and repeat subtotal resection because of base-of-tongue and epiglottis involvement. A postoperative swallow study revealed normal function without aspiration. A definitive serial CO₂ laser excision was then performed to remove all residual disease, and the patient was successfully decannulated. At last follow-up approximately 1 year later, the patient continued to be free of disease.
Discussion

Myofibromas are benign neoplasms composed of contractile myoid cells and myofibroblasts; they are considered the most common fibrous proliferation of infancy and childhood.\(^1,2\) Although an estimated 90% of all cases occur in the first 2 years of life,\(^3\) myofibromas can arise in patients of any age and, accordingly, the condition is broadly divided into infantile (juvenile) myofibromatosis and adult myofibromatosis. This lesion can manifest as solitary or multicentric, and it is believed that solitary lesions are more common.\(^4\)

Myofibromas have been reported to have a predisposition for the subcutaneous and cutaneous tissues and skeletal muscles of the head and neck—in particular the oral cavity.\(^3,4\) Here we report the first case of a solitary oropharyngeal myofibroma in an adult and describe its potential for life-threatening airway compromise.

Myofibromas are generally considered to be slowly growing tumors most frequently seen on the oral tongue and buccal mucosa. When present, these tumors can be difficult to distinguish from other fibrous proliferations, including fibrosarcomas, leiomyomas, fibrous histiocytomas, and solitary fibrous tumors, among others. Reports of the histopathologic features of solitary myofibromas have varied, ranging from a strictly benign appearance to a locally aggressive mass with infiltrative growth and high rates of recurrence after excision.\(^4\)

In our case, the lesion was noted to be a bland, spindle cell neoplasm with cells exhibiting elongated, cigar-shaped nuclei with normochromasia and without mitoses or necrosis. Immunohistochemistry revealed cytoplasmic positivity for smooth-muscle actin, seen in myofibroblasts, and negative staining for several keratins (positive in spindle cell carcinomas), CD34 (positive in solitary fibrous tumors), ALK1 (as seen in inflammatory myofibroblastic tumors, usually pediatric), and desmin (positive in leiomyomas). The overall findings were of a benign myofibroblastic proliferation consistent with myofibroma.

Once the diagnosis of myofibroma is established, complete excision with a conservative surgical resection has been advocated, especially if the tumor is in a functionally or cosmetically sensitive location.\(^2\) In the present case, epiglottic involvement necessitated initial subtotal resection to minimize the risk of subsequent aspiration while the airway was stabilized, followed by definitive laser excision.

Myofibromas also have been reported to be successfully treated with serial subtotal resections followed by complete spontaneous regression, which is a phenom-
patient who required 4 days of oral supplementation before normalization.

One shortcoming of our study, in addition to its retrospective nature, is that intact parathyroid hormone levels had not been obtained in the hypocalcemic patients. The availability of these measurements would have allowed us to refine our assessment.

In conclusion, our findings suggest that minimally invasive thyroidectomy poses no additional risk with regard to the development of prolonged postoperative hypocalcemia. We did not seek to compare the minimally invasive thyroidectomy cohort with our patients who had undergone conventional open thyroidectomy because they were dissimilar groups with respect to thyroid anatomy, disease state, and the extent of resection. These differences made any such comparison of little value in terms of the goal of this study.

References

many occasions and should be considered in the differential diagnosis of any subcutaneous or submucosal head and neck lesion. This case should alert clinicians to the potential obstructive nature of these lesions when present in solitary or multicentric forms in both children and adults. Management principles include airway stabilization, surgical excision with preservation of speech and swallow function, and close postoperative monitoring for recurrence.

References