A Giant Combined Laryngomucocele

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ABSTRACT

Laryngomucocele is a rare benign laryngeal disease characterized by cystic dilatation of laryngeal saccule with collection of mucous in it. The etiology behind its occurrence is still unclear, but congenital and acquired factors have been implicated in its development. Although it is a benign lesion, its presence in an old patient since many days should prompt the examiner to diligently search for laryngeal malignancy. We are presenting the case of a 61 years old male who presented with swelling in the neck, hoarseness of voice, dysphagia and dyspnea on exertion. The patient was thoroughly evaluated clinically, endoscopically and radiologically. He was diagnosed to be suffering from laryngomucocele and was managed by an external lateral cervical approach.

Keywords: Laryngomucocele, Laryngocele, Larynx.

INTRODUCTION

Laryngocele is defined as an abnormal dilatation or herniation of the saccule of the larynx. Laryngoceles can be classified into three types: internal, external and combined. The correlation of laryngocele and laryngeal carcinoma is still being debated. Laryngocele may be seen in certain professions where high intraglottic pressure needs to be maintained for a long period of time, but it is not necessary that such a history be present in all cases. Patient presents generally with hoarseness of voice and swelling in the neck. Definitive diagnosis is made by computed tomography (CT) or magnetic resonance imaging (MRI) scans. Treatment is endoscopic marsupialization for internal laryngocele and external approach for external or recurrent laryngoceles.

CASE REPORT

A 61 years old male presented in ENT OPD with a history of progressive swelling in the right side of neck and hoarseness of voice since 3 years. He had difficulty in swallowing solids and dyspnea on exertion from 6 months. He gave a history of the swelling, getting aggravated on doing strenuous activity. The swelling which was initially reducible, became nonreducible from the past one and a half years. He is a farmer by occupation and used to sing devotional songs aloud.

On examination, there was a swelling over the right side of neck medial to anterior border of sternocleidomastoid with approximately 6 × 4 cm in size extending vertically from hyoid bone to cricoid cartilage. The swelling was cystic, nontender and compressible but nonreducible. The swelling became prominent on performing Valsalva maneuver (Fig. 1).

Indirect laryngoscopy showed bulge at right vallecula pushing the epiglottis and laryngeal inlet toward the left, right pyriform sinus could not be seen (Fig. 2). Seventy degree endoscopy was done to confirm our findings. Ultrasound of the neck revealed cystic swelling in the right paralaryngeal region with dehiscence in the thyrohyoid membrane. Thick fluid was aspirated on fine needle aspiration cytology (FNAC) which showed no inflammatory or malignant cells. Computed tomography scan of the neck revealed a large rim enhancing dumb-bell shaped cystic lesion on right side of neck at the level of larynx with superficial and deep components, possibly a mixed fluid filled laryngocele (Fig. 3). A presumptive diagnosis of mixed laryngomucocele
was made and the patient was managed by doing complete surgical excision by external lateral cervical approach. The external and internal components were communicating through a defect in the thyrohyoid membrane. Right-sided superior laryngeal nerve was identified and preserved. A triangular wedge of thyroid cartilage was resected in order to gain accesses to the internal part. The laryngomucocele was removed in toto and the neck of saccule was ligated. There were no complications during or after surgery. Histopathology of the specimen revealed cyst wall lined with respiratory epithelium, and there was no evidence of any malignancy. The patient is under follow-up from the past one and a half years and has been asymptomatic till date (Fig. 4).

DISCUSSION

Laryngocele was first described by Virchow in 1863. He used the term ‘laryngocele ventricularis’ to describe an anomalous air sac, communicating with the laryngeal ventricle. Upward extension from the anterior end of the laryngeal ventricle is the laryngeal saccule. Saccule is a pouch arising from the anterior end of laryngeal ventricle. The laryngeal ventricle is the space between the vocal and vestibular folds. Saccule extends superiorly between false vocal fold and the inner surface of thyroid cartilage. A laryngocele is an abnormal dilatation or herniation of the saccule that communicates with the lumen of the larynx, and is air filled. If the communication between the laryngocele and the laryngeal lumen gets occluded then fluid gets accumulated within the sac. If the accumulated fluid is mucoid in nature the term ‘laryngomucocele’ is used or if it gets secondarily infected and pus gets accumulated then ‘laryngopyocele’ is used to define the mass.

Laryngocele has a male predominance with male to female ratio of 5:1 and peak incidence is in the sixth decade of life.2 Laryngoceles may be congenital or acquired. Although the etiology remains unclear but few propose that the cause of acquired laryngocele may be a significant increase in intralaryngeal pressure for a long period as in glass blowers, wind instrument players and singers. Chronic cough could be one of the precipitating factors. Certain conditions can cause the closure of the neck of saccule leading to laryngocele formation by a flap valve mechanism, i.e. allowing entry of air in the saccule but preventing its egress like neoplasm associated with ventricle and false cord, inflammatory conditions and sometimes trauma.3,4

Three types of laryngoceles have been described. Internal laryngocele is one which is confined to the interior of the larynx and extends posterosuperiorly into the false cord and the aryepiglottic fold.5 If the laryngocele extends superiorly and appears laterally in neck through an opening in the thyrohyoid membrane, this is external laryngocele.6 Combined type has an external swelling in the thyrohyoid region which expands on coughing and empties on digital pressure and an internal component seen as a smooth

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**Fig. 2:** Preoperative endoscopic view showing internal component of laryngomucocele

**Fig. 3:** Computed tomography scan of the neck revealed a large rim enhancing dumb-bell-shaped cystic lesion on the right side of the neck

**Fig. 4:** Postoperative endoscopic view showing normally aligned larynx
submucosal swelling in the vallecula and obliterating the pharyngoepliglottic fold.

Combined laryngocele appears to be the most common type of laryngoceles as compared to external or internal types. Classically, the swelling will increase during a valsalva maneuver and become smaller on palpation. Internal and combined laryngoceles appear on laryngoscopy as a smooth swelling of supraglottis.

Laryngocele may be asymptomatic and often incidentally discovered through radiographic studies for unrelated diseases. Patient presents mainly with airway obstruction, stridor, hoarseness, sore throat, cough, pain or an externally palpable mass. It has been documented that laryngomucocele are associated with laryngeal carcinoma. External laryngocele has been found in 16% of laryngectomized specimen operated for laryngeal carcinoma. Patients with external or mixed laryngocele may present with a swelling on one or both sides of the neck, which is usually compressible and increase in size on performing valsalva maneuver. There is no predominance of occurrence on left or right side.

The diagnosis can be reached by assessing the patient clinically, by doing a laryngoscopy (direct/fiberoptic) and imaging studies including soft tissue radiograph and CT of the neck and MRI. Magnetic resonance imaging appears now to be the modality of choice due to better soft tissue resolution.

Asymptomatic lesion can be managed conservatively. Symptomatic lesions have to undergo surgical excision. Cosmetic disfigurement is a relative indication for excision. Management depends upon the size of the lesion. Laryngoceles have traditionally been treated through an external approach. Myssiorek et al have used lateral laryngectomy approach for resection of laryngoceles and stated that the external approach without tracheostomy allowed for good exposure with minimal functional disability. Small internal laryngoceles can be approached using a CO₂ laser endoscopically or by an external incision. External or combined laryngoceles need an external cervical approach. In combined laryngocele, removal of a triangular wedge of thyroid lamina gives a wide exposure to paraglottic space, thus helping in dissection of the internal component of the laryngocele. The external lateral cervical approach hardly causes any functional disability. During the procedure, the superior laryngeal nerve must be identified and carefully preserved as it could be intimately related to the mass.

CONCLUSION

Laryngomucocele is a rare benign laryngeal disease which is often asymptomatic. The present case is of interest because the patient was affected by a very large nonreducible external swelling, presented with exertional dyspnea, hoarseness of voice and dysphagia. The present case was thoroughly investigated with an interest to rule out associated laryngeal cancer which is a common cause for laryngomucocele in an elderly person. An external lateral cervical approach to the laryngomucocele gave an adequate exposure and postoperative recovery was free of complications. There was no recurrence on follow-up for one and half years.

REFERENCES