Left mixed laryngocele in absence of risk factors: A case report and review of literature

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ABSTRACT

Laryngocele is an uncommon benign cystic dilatation of the laryngeal saccule that communicates with the laryngeal lumen and contains air. On the basis of its localization, it can be traditionally classified in internal, external, or mixed. Usually unilateral and rarely bilateral, it may be congenital or acquired. It most often appears in severe cases; thus, it is termed laryngopyocele [1]. Accordingly to the traditional classification, laryngocele can be described as internal (within the larynx), external (outside the larynx), or mixed (both) [1]; however, many authors currently prefer to reduce this classification to only internal and mixed laryngocele [3]. Usually unilateral, laryngocele recognizes several etiologies including congenital, increased laryngeal pressure, and mechanical obstruction; it is often asymptomatic and found incidentally on imaging [2]. The treatment of choice is surgical excision [1,3], microlaryngoscopic, external or combined, depending on the laryngocele characteristics.

Here, in line with the SCARE criteria [4], we describe the case of a 72-year-old man smoker but without other specific risk factors, who presented laryngeal dyspnea for about one year during hospitalization in another department for respiratory failure. In the context of this hospitalization, a neck CT-scan was performed revealing a left mixed laryngocele (Fig. 1). The patient was a farmer and currently retired; he was a smoker but had a negative history of alcohol consumption. He denied other risk factors for laryngocele. He suffered from high blood pressure and COPD, but had a negative history of alcohol consumption. He denied other risk factors for laryngocele. He suffered from high blood pressure and COPD, had already undergone gastroresection for gastric ulcer about 30 years ago and at time of clinic evaluation he denied other symptoms besides dyspnea.

On examination, he presented a left submandibular mobile mass not painful on palpation that was covered with normal skin (Fig. 2). A 72-year-old man presented to our attention with a history of laryngeal dyspnea for about one year during hospitalization in another department for respiratory failure. In the context of this hospitalization, a neck CT-scan was performed revealing a left mixed laryngocele (Fig. 1). The patient was a farmer and currently retired; he was a smoker but had a negative history of alcohol consumption. He denied other risk factors for laryngocele. He suffered from high blood pressure and COPD, had already undergone gastroresection for gastric ulcer about 30 years ago and at time of clinic evaluation he denied other symptoms besides dyspnea.

On examination, he presented a left submandibular mobile mass not painful on palpation that was covered with normal skin (Fig. 2). Video rhinofibrolaryngoscopy showed a smooth-walled cystic swelling

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occupying the left portion of the supraglottic larynx, from the ariepiglottic fold to the ventricular folds and ventricle of Morgagni. A neck MRI was performed to confirm the diagnosis and better establish the boundaries of the mass: it showed how the laryngocele reached the infero-medial pole of the left submandibular gland causing an extrinsic compression of laryngeal vestibule and a right shift of the glottic lumen (Fig. 3).

Three days after admission to the otolaryngology department, after adequate written informed consent, surgery was used to excise the laryngocele via cervicotomic access under general anesthesia. The first time consisted of tracheotomy under local anesthesia in order to facilitate the next phase; subsequently, the left submandibular region over the swelling was incised under general anesthesia, dissecting until reaching the thyrohyoid membrane where the laryngocele was isolated and removed entirely (Fig. 4).

The subsequent post-operative course was regular and free of complications. The tracheal cannula was removed on day three. On the 7th post-operative day, the patient was discharged in good health and with final histological diagnosis of laryngocele. One month after surgery, the patient attended an outpatient hospital check-up visit in the context of which videolaryngoscopy established the absence of disease (Fig. 5).

3. Discussion

Laryngocele was first described by Larrey (a surgeon in Napoleon’s army) in 1829 as “an air-filled tumor” and then by Virchow who gave it the name by which it is known today in 1887 [1]. It is an abnormal cystic dilation of laryngeal saccule [1,2], a small mucosal pouch situated in the upper part of ventricle of Morgagni between the ventricular folds, the base of epiglottis, and the inner surface of thyroid cartilage [1].

Being in communication with laryngeal lumen, laryngocele normally contains air, but by virtue of the presence of mucinous glands in the ventricle, it may contain mucus (laryngomucocele) or, in severe cases of infection, it may be filled with pus (laryngopyocele) [2].

With an incidence of 1 per 2.5 million people per year, laryngocele is a rare clinical entity, five times more common in men than women [5]. It may present at any age from neonates to the elderly, but the reported peak incidence in the literature is between the fifth and sixth decade of life [1,5]. It is unilateral in about 85% of cases [6] and in more than 8% of cases it turns into a laryngopyocele [7], that is the main complication.

Laryngocele is historically classified as internal when inside the larynx, confined within the thyrohyoid membrane; external if outside the larynx, beyond the thyrohyoid membrane and extended to the subcutaneous tissues of the neck; mixed if it presents both components [2,8]. However, many modern authors prefer to reduce this classification only to internal and mixed laryngocele [2]: therefore, those laryngoceles medial to the thyrohyoid membrane are internal, and mixed those located beyond the membrane [9]. The latter are also the most common type of laryngocele, making up 50% of all [7].

Regarding its etiologies, laryngocele can be both congenital and acquired. It is most often congenital and related to a large ventricle appendix [10]. Among acquired causes, perhaps the most important are all those conditions that lead to an increased intraluminal laryngeal pressure for repeated and prolonged periods: thus, laryngocele is more common in glass blowers and wind instruments players [8]. From this point of view, also mechanical obstructions of the larynx that increase the endoluminal pressure can lead to laryngocele as in the case of laryngeal obstructive carcinoma, especially supraglottic, amyloidosis, chondroma, rheumatoid arthritis, or scleroderma [1,3]. However, adult patients at highest risk for development of laryngocele are usually those with congenitally dilated saccules [10]: in fact, in them also the Valsalva maneuver, that accosts true and false vocal folds increasing intra-laryngeal pressure, might exacerbate the congenitally dilated sacculle to result in a laryngocele [2]. Nevertheless, all these triggering conditions were absent in the patient subject of case report, for which it was not possible to recognize any known risk factors.

The diagnosis of laryngocele is essentially clinical: its most common symptoms are hoarseness and neck swelling, but it may also present with snoring, upper airway obstruction, increasing stridor, sore throat, cough, pain, globus sensation, and dysphagia [1,3,11]. In our case, patient presented dyspnea and acute respiratory failure like pulmonary pathology. Nevertheless, laryngoceles are very often asymptomatic and therefore diagnosed incidentally with imaging [8] especially CT, which
is currently the most accurate imaging modality for their diagnosis [2,8] and in which laryngocele appears as an air-filled structure in the space near the laryngeal sacculle. Another diagnostic possibility is MRI, which is useful to define boundaries of the dilatation and its relationship with thyrohyoid membrane, distinguishing internal from external and mixed laryngocele [1]; moreover, MRI can discriminate laryngocele from laryngomucocele, laryngopyocele, and even the eventual underlying laryngeal squamous cell carcinoma [10].

As previously mentioned, the main complication of laryngocele is pyocele, but other complications also include pathogens aspiration with subsequent bronchitis and pneumonia, rupture and following infection in the lateral larynx, and upper airway obstruction [6]; for this last reason, both laryngoceles and laryngopyoceles are possible causes of sudden death [7].

The treatment of choice is surgical excision [1,3] for which different approaches are possible depending on the characteristics of the lesion. For example, smaller internal laryngoceles are managed via micro-laryngoscopy with a CO₂ laser [1,9], which presents fewer complications and faster post-operative hospitalization [9], but with the disadvantages of limited surgical exposure, post-operative endolaryngeal scarring, and risk of incomplete resection [2,9]. On the contrary, larger or mixed laryngoceles are approached through external access on the lateral neck [1,2]. This last method is preferred for superior exposure, precision, and a low recurrence rate; however, it implies higher morbidity, more complications (edema and subsequent airway impairment, laryngocutaneous fistula, subcutaneous emphysema and nerve injury), longer surgery, and longer recovery time with subsequently increased cost [2]. The external approaches described in the contemporary literature include the transthyrohyoid membrane approach (most common), thyrotomy with resection of the upper 1/3 of thyroid cartilage, and V-shaped thyrotomy. The latter two both offer a superior visualization of the paraglottic space relative to the transthyrohyoid membrane approach [2,9]. To prevent the risk of respiratory distress, a tracheotomy is practiced as a surgical first step in many patients [1], as in our case.
In conclusion, laryngocele is a rare condition that recognizes an increased intraluminal laryngeal pressure as main risk factor, absent in our patient. Although its main symptoms, if present, are dysphonia and/or dysphagia, it may also manifest in atypical way, with an acute respiratory failure simulating a pulmonary pathology, as reported in this clinical case. Nevertheless, laryngocele is an asymptomatic condition in most cases, discovered incidentally on imaging, especially on CT, and whose main treatment is surgical through external approaches for mixed laryngoceles and microendoscopic for internal ones.

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Ethical approval

For all procedures performed in case report “Left mixed laryngocele in absence of risk factors: a case report and review of literature”, formal consent of institution’s ethics committee is not required.

Registration of research studies

1. Name of the registry:
2. Unique Identifying number or registration ID:
3. Hyperlink to your specific registration (must be publicly accessible and will be checked):

Guarantor

Bisogno Antonella.

Declaration of competing interest

The authors declare no conflict of interest.

Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.amsu.2020.11.024.

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Author contribution

Bisogno Antonella: data collection; literature review; writing the paper.
Cavaliere Matteo: data analysis and interpretation; correction of the manuscript; surgeon.
Scarpa Alfonso: data analysis and interpretation; correction of the manuscript; surgeon.
Cuofano Rossella: data analysis and interpretation; surgeon.
Troisi Donato: data collection; literature review; surgeon.
Iemma Maurizio: study concept and design; first surgeon; director of Otorhinolaryngology Department.