Cavernous lymphangioma in the laryngeal ventricle in an adult patient

Erişkin bir olguda larenks ventrikülünden kaynaklanan kavernöz lenfanjiom

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Lymphangiomas of the larynx are quite rare and even rarer in adults. Isolated cavernous lymphangioma in the laryngeal ventricle was diagnosed in a 47-year-old woman. The patient presented with hoarseness with a history of three months. Direct laryngoscopy performed under general anesthesia showed a large, smooth, gray-reddish tumor above the right vocal fold, originating from the right ventricle. The tumor was removed through a micro-laryngoscopic procedure with cold instruments. Histopathologic diagnosis was cavernous lymphangioma. There was no local recurrence a year after the operation.

Key Words: Adult; hoarseness/etiology; laryngeal neoplasms/complications; lymphangioma/pathology/surgery.


Anahtar Sözcükler: Erişkin; ses kısıklığı/etioloji; larenks neopazileri/komplikasyon; lenfanjiom/patoloji/cerrahi.

Lymphangiomas are rare congenital lesions of the lymphatic system, most commonly presenting in the head and neck during infancy. Of all lymphatic malformations, 50% to 60% are present at birth, and approximately 80% to 90% become evident by the age of two years. The occurrence is uncommon in the older child, and even rarer in the adulthood. Lymphangioma of the larynx is also an extremely rare incidence. Naito et al. reviewed the entire English literature and documented 17 lymphangioma cases with isolated laryngeal involvement in adults and children. Only four adult cases appeared in the English literature within 2002.

Kennedy classified lymphangiomas into four types: superficial cutaneous, cavernous, cystic hygroma, and diffuse systemic. Cavernous lymphangiomas are composed of dilated lymphatic channels with one or more endothelial layers with or without an adventitial layer.

We present an adult patient with a cavernous lymphangioma confined to the laryngeal ventricle.
CASE REPORT

A forty-seven-year-old female patient presented with hoarseness with a history of three months. Indirect laryngeal examination revealed a large, smooth, gray-reddish mass above the right vocal fold. Her breathing was normal. Direct laryngoscopy under general anesthesia showed a large tumor originating from the right ventricle (Fig. 1). The right true vocal cord was poorly visualized because of the extension of the mass along the right laryngeal ventricle. A red, smooth mass, 11x9x5 mm in size was removed by way of a micro-laryngoscopic procedure with cold instruments. The remaining parts of the larynx appeared normal. She was discharged on the day after surgery.

Histopathologic examination showed spaces of dilated, irregular shape and size under the thick stratum of ciliated columnar epithelium, lined by the endothelium. Some of them contained a small number of lymphocytes and a pinkish pale coagulated amorphous mass, leaving the greater parts empty. The stroma was composed chiefly of fibroblasts and focal lymphocytic aggregates near the vessels (Fig. 2, 3). The diagnosis was made as cavernous lymphangioma.

The complaints of the patient completely disappeared; her voice became full-sounding and clear. No recurrences were detected a year after surgery.

DISCUSSION

Lymphangioma of the larynx mainly occurs in regions where friction is caused by the passage of food and where there are numerous lymphatic vessels, such as the lingual surface of the epiglottis, aryepiglottic folds, and the arytenoid region. Its appearance is very rare in the vocal folds, in the ventricles of Morgagni, and in the subglottic area. Lymphangiomas do not infiltrate the surrounding

Fig. 1 - Micro-laryngoscopic view (7x) showing a large, gray-reddish, smooth mass of the right vestibular fold. The mass was fairly solid to palpation. It was confined to the right laryngeal ventricle.

Fig. 2 - Under the thick stratum of ciliated columnar epithelium, there were spaces of dilated, irregular shape and size, lined by the endothelium. Some of them contained a small number of lymphocytes and a pinkish pale coagulated amorphous mass, but the greater parts were empty (H-E x 40).
tissues because of a distinct capsule. Those with a small size and location other than the vocal folds usually cause few or no symptoms. However, their growth or location in the sphere of the glottis may be associated with changes in voice and may lead to hoarseness, cough, nausea, vomiting, dispnea, and disphagia.\[6\]

Lymphangioma rarely occurs in the larynges of infants and children. Its appearance is uncommon in the older child, and even rarer in the adult. Compared to that of the adults, the lymphatic system of the larynx is better developed in infants and young children, with a greater number of channels and lymph nodes.\[8\] Cohen and Thompson\[8\] reviewed 160 pediatric patients with lymphangiomas, of whom ten had extensive laryngeal involvement accompanied by cervical, facial, and/or submandibular hygromas.

To our knowledge, there are few literature reviews giving detailed information on laryngeal lymphangioma in adults. Gerwel and Nawrocki\[1\] reported a 17-year-old girl with complaints of difficulty in swallowing, change in voice, and a feeling of a foreign body in the pharynx. Laryngeal examination revealed a large tumor involving the hypopharynx and the whole lingual surface of the epiglottis, which was then removed via suprathyroid pharyngotomy together with the epiglottis at the base of the tongue. Smith and Stafford\[10\] reported a 69-year-old female patient with lymphangioma of the post-cricoid region. She presented with symptoms characteristic of globus pharyngeus. It was removed via pharyngoscopy. Naito et al.\[12\] reported a 36-year-old female patient who complained of hoarseness for several months associated with lymphangioma on her right false vocal cord. They performed tracheostomy and a laryngofissure for the removal of the neoplasm. We could not find any adult case with lymphangioma in the laryngeal ventricle in the English literature.

The recurrence of lymphangiomas ranges from 39% to 50% in pediatric patients following surgery.\[5,6\] The recurrence was 21% after attempted total resection as the initial procedure.\[7\] In our case, no recurrence was detected within a year.

REFERENCES

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