Complete Congenital Third Branchial Fistula: A Rare Case Report

Sujata A Gawai, Kalpana R Kumar, Vaishali S Sangole, Suman P Rao, Divya A George, Rachna Tiwari, Bhagirath D Kandhare

1 Senior Resident, Department of ENT, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India
2 Professor, Department of ENT, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India
3 Lecturer, Department of ENT, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India
4 Professor and Head, Department of ENT, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India
5 Resident, Department of ENT, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India
6 Assistant Professor, Department of Radiology, MGM Medical College and Hospital, Kamothe, Navi Mumbai, Maharashtra, India

Correspondence: Sujata A Gawai, Senior Resident, Department of ENT, MGM Medical College and Hospital, Sector 18, Kamothe Navi Mumbai-410209, Maharashtra, India, Phone: +919987361256, e-mail: drsujatagawai@gmail.com

ABSTRACT

Fistulae arising from the second branchial apparatus are the most common anomalies than those arising from the third and fourth branchial apparatus. Third branchial fistula is extremely rare in occurrence. Complete fistulae are uncommon as in the majority of cases the tracts end blindly. Here, we report a case of complete congenital third branchial fistula on left side of the neck with an internal opening near left pyriform fossa in a 12-year-old male patient.

Keywords: Branchial cleft, Fistula.

INTRODUCTION

More than 90% of branchial cleft anomalies arise from second branchial cleft system and 8% from the first branchial cleft system. Anomalies arising from third and fourth branchial cleft system rarely occur. Persistence of remnants of the branchial apparatus gives rise to a number of well-recognized congenital anomalies in the head and neck. A persistent cleft will give rise to an external sinus, a blind ending opening onto the skin. A persistent pouch will cause an internal sinus typically opening into the pharynx, whereas persistence of both cleft and pouch will cause a fistula with an internal and external opening. Even if an internal opening exists, there lies a thin mesodermal tissue between the external and internal opening. A complete branchial fistula develops from a rupture of membrane between the cleft and pouch at the same time during development. Treatment requires complete excision of the tract in order to prevent recurrence.

CASE REPORT

A 12-year-old male patient presented to ENT OPD of MGM hospital with history of discharge from the left side of neck since the age of 4 months. He had a history of swelling in the left side of neck at 4 months of age which bursted spontaneously with mucoid discharge through a small opening at the same site. Ever since then there was watery discharge from the opening while drinking.

Examination revealed a small pinpoint opening in the left side of the neck over the anterior border of left sternocleidomastoid muscle, 2 cm above the medial end of left clavicle (Fig. 1A). Surrounding the opening there was a

Fig. 1A: External opening of the branchial fistula at the anterior border of left sternocleidomastoid muscle, 2 cm above medial end of left clavicle

Fig. 1B: Watery discharge seen after giving sip of water to drink
hyperpigmented area with an old healed scar. Watery discharge was seen after giving a sip of water to the patient for drinking (Fig. 1B).

General examination was within normal limits.

X-ray barium swallow was performed to know the site of internal opening but it failed to show any tract. X-ray fistulogram was done by cannulating the opening and by injecting diatrizoate meglumine and diatrizoate sodium 76% after diluting, a fistulous tract was revealed extending superomedially to open in the left side of hypopharynx (Fig. 2).

MRI was performed to know the course of the tract and the site of internal opening, which showed a fistulous communication of the cutaneous opening with left pyriform fossa (Figs 3A and B). The track was found anterosuperomedial to the left carotid sheath.

For exploration and excision of the fistula, patient was taken under general anesthesia, the external opening was canulated with 24 number angiocatheter, and methylene blue dye was injected which spilled into the pyriform sinus and revealed the internal opening on direct laryngoscopy. A transverse elliptical incision was made around the external opening and subplatysmal flaps were raised carefully, the

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**Fig. 2:** X-ray fistulogram of branchial fistula

**Fig. 3A:** MRI of neck shows tract running anterosuperomedial to the left carotid sheath

**Fig. 3B:** MRI of neck shows fistulous tract opening in left pyriform fossa

**Fig. 4:** Intraoperative picture of branchial fistula being excised

**Fig. 5:** Histopathological picture showing tract lined by keratinized stratified squamous epithelium
tract was identified as firm cord-like structure, which was gradually dissected from the surrounding tissues. The tract was found anteromedial to the carotid sheath and was not going deep to the carotid arteries as described in the literature. Aneeza and Mazita et al reported in 2010, a complete third branchial fistula which was also anteromedial to the carotid sheath.6

Symptoms consist of intermittent or continuous discharge and recurrent attacks of inflammation following an attack of upper respiratory tract infection. The external opening may be seen to move upward with deglutition.9 For investigating the complete branchial fistula, barium contrast study or fistulogram can be performed which will clearly delineate the tract. In difficult cases both CT scan and MRI can be done.

Surgery is indicated to prevent the risk of recurrent infection and for cosmetic reasons. Treatment is complete surgical excision of the tract, in order to avoid its recurrence. Different methods for surgical excision of the tract have been used, the standard method includes stepladder excision for extensive fistulas. In our case, single incision around the external opening was sufficient to excise the tract completely. Another method is stripping the tract using wire stilletes, vein strippers or arterial intimal strippers; however, being a blind procedure, it is not generally advocated if vital structures exist in the proximity.10 The recurrence rate of branchial anomaly is 3% for a primary lesion and as high as 22% for lesions with previous infection and surgery.11 Complications of the surgery include secondary infection, injury to facial, hypoglossal, glossopharyngeal, spinal accessory nerves, injury to internal jugular vein, bad scar and hematoma formation.12

Endoscopic cauterization of the fourth branchial cleft sinus has been reported with effective result compared with open excision.13 Treatment by chemocauterization of the internal opening with trichloroacetic acid (TCA) has been reported with encouraging results.2,14

In 2009, Nixon and Healey reported the role of sclerotherapy for treatment of branchial sinus tract. They used 3% sodium tetradecyl sulphate foam delivered via catheter along the length of the tract with significant improvement in the fluid discharge.15

In this article, we discussed a case of complete congenital third branchial fistula, which is rare in occurrence. The presence of a complete tract and its diagnosis by careful and complete history, physical examination and necessary investigations including fistulogram is very important for the complete treatment of the fistula.

REFERENCES


Editorial Inputs

Complete Third Branchial Arch Fistula: A Rare Presentation (Multiple Surgery Failure)

Saurabh Varshney
Professor and Head, Department of ENT, Himalayan Institute of Medical Sciences, Dehradun, Uttarakhand, India

The incomplete branchial fistula is not an uncommon congenital anomaly of branchial apparatus, but a complete one is rare. Anomalies arising from the third and fourth branchial cleft system rarely occur. Third and fourth branchial pouch fistulas are collectively referred to as pyriform sinus fistulas. Branchial cleft sinuses with external openings are usually associated with the first and second branchial cleft arches, and those with internal openings are usually associated with the third and fourth arches. The course of a third branchial fistula is derived from its embryological origin, in accordance with the branchial apparatus theory.
We have treated 3 such cases in last 5 years at our institute, the last case treated by us was a 15-year-old male patient who presented with a history of a small opening with mucus discharge from the right side of the neck since birth, which increased on taking food. He had no family history of similar complaints. There was history of surgical excision of the fistula tract 9 times under general anesthesia without cure. Computed tomography with contrast injection into the external cervical opening revealed a patent tract from the neck skin to the base of the pyriform sinus, which was also confirmed by injecting methylene blue from external opening and viewing it in right pyriform sinus with laryngeal endoscope. Complete excision of the tract up to the pyriform sinus with right hemithyroidectomy and closure of defect in right pyriform sinus was performed. Follow-up at 6 months has no recurrence. Our case with multiple surgery failures emphasizes that treatment of this condition requires complete removal of the tract in order to avoid recurrence; however, this can pose a risk to the surrounding structures, hence many surgeons perform incomplete excision of tract resulting in recurrence.

### Editorial Inputs

**Gauri Mankekar**

Branchial arch fistulas are easy to diagnose but difficult to treat. Often the tract vanishes between the carotid bifurcation and one cannot trace its entire length. Patients often undergo multiple surgeries before they are free of the discharging fistula.