Case Report

First branchial cleft fistula: a case report

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ABSTRACT

First branchial cleft anomalies (BCA) are a rare finding in head and neck with incidence of nearly 1 million per year which are distributed below external auditory canal, above the hyoid bone, anterior to sternocleidomastoid and posterior to submandibular triangle. First branchial cleft cysts are frequently misdiagnosed as they are rare and pose unfamiliar clinical signs and symptoms. Here we are reporting a case of surgical management of 1st branchial cleft fistula in a 5 years old male child from AIIMS, Raipur, Chhattisgarh, India as it’s a rare entity. Child presented with discharge from right upper part of neck. There was a swelling in right upper lateral part of neck with an opening also in floor of right external auditory canal (EAC). Contrast enhanced computed tomography of neck showed a 4.8 cm long obliquely oriented fistulous tract opening at junction of middle and upper one third of sternocleidomastoid with opening in right EAC. Surgical excision of the fistulous tract was done with preservation of facial nerve. Histopathology examination confirmed the presence of fistula. Common clinical presentation of BCAs is pre-auricular swelling (24%), parotid swelling (36%) or cervical region swelling (41%). In our case, it was a fistulous opening that presented as discharging tract in upper neck. Management include early diagnosis, control of infection and complete excision with facial nerve preservation Surgical approach should be based on the clinical examination, imaging and clinical course; and there is a need to safely identify and preserve facial nerve in almost all cases.

Keywords: Branchial cleft anomalies, Facial nerve, Branchial cleft fistula

INTRODUCTION

The term ‘branchial cyst’ was coined by Ascherson in in 1832.1 First branchial cleft anomalies are a rare finding in head and neck, the incidence being around 1 million per year.1,3 The occurrence of first branchial cleft anomalies usually varies from less than 8% to 21.4%.1,3 They are distributed below external auditory canal, above the hyoid bone, anterior to sternocleidomastoid and posterior to submandibular triangle.1 Main theories which describe the development of branchial cleft anomalies include incomplete obliteration of branchial mucosa, persistence of vestiges of pre-cervical sinus, origin from thyropharyngeal duct and origin from cystic lymph nodes.1

In 1972, Work proposed a classification for first branchial cleft anomalies which states that type 1: anomalies superficial to facial nerve in close proximity to ear and type 2: lesion communicating with external auditory canal or tympanic membrane, mostly lying medial to facial nerve.3

First branchial cleft cysts are frequently misdiagnosed as they are rare and pose unfamiliar clinical signs and symptoms.4 Management of these includes early diagnosis, control of infection and complete excision with facial nerve preservation.3 Here we present a case of first
branchial cleft fistula in a 5 years old male child and its surgical management.

CASE REPORT

A 5-year-old male child visited department of Otorhinolaryngology and Head and Neck Surgery in outpatient department at AIIMS Raipur with complaints of recurrent discharge from right upper neck since last 3 years. According to his parents, the child had 2 to 3 such episodes per year on an average which began as painless swelling which later ruptured spontaneously. On various occasions it was associated with ear discharge which subsequently resolved with medication. Child had unremarkable family and medical history. There was a swelling at the junction of upper and middle one third of sternocleidomastoid muscle at its anterior border with small opening in the floor of right external auditory canal with no active discharge and no cervical lymphadenopathy.

Management and surgical approach

USG neck revealed a cutaneous lesion in right paramedian region in anterior cervical triangle with irregular hypoechoic sinus tract, suggestive of branchial cleft sinus. Contrast enhanced computed tomography of neck showed a 4.8 cm long obliquely oriented fistulous tract opening at junction of middle and upper one third of sternocleidomastoid, then extending superiorly and laterally along the anterior border of sternocleidomastoid travelling behind right submandibular gland posterior to ramus of mandible and deep to parotid gland opening internally at anteroinferior and medial aspect of right auricle.

Based on the investigations, diagnosis of first branchial cleft fistula was made and excision under general anaesthesia was planned.

(Figure 1). Methylene blue dye was injected through the opening in upper neck.

![Figure 1: Incision with lower elliptical extension around the opening in lower part of neck.](image1)

Modified Blair’s incision was given behind and below the right pinna and extended below till the opening in the upper neck which was elliptically included in the incision.

![Figure 2: Exposure of the fistulous tract.](image2)

![Figure 3: Dissection of the fistulous tract preserving the facial nerve.](image3)

![Figure 4: Excised fistulous tract.](image4)
Dissection was started from lower elliptical end which was dissected from surrounding tissue and raised upwards along the anterior border of sternocleidomastoid, posterior to submandibular salivary gland, further traversing medial to parotid gland and facial nerve trunk (Figures 2-5).

With preservation of all the aforementioned structures, upper end of fistula was traced which was just lateral to right external auditory canal in the auricle. Entire fistula tract, nearly 6.5 cm in length was excised and sent for histopathological evaluation. Patient was discharged on postoperative day 4, intravenous antibiotics were administered. Histopathological analysis revealed skin with underlying tissue containing fistulous tract lined by stratified squamous epithelium with adnexa.

Oral antibiotics were advised and patient was called for follow up on post-operative day 7 for suture removal. On follow up, surgical site was healthy. Further monthly follow-up was advised.

DISCUSSION

First branchial cleft anomalies can appear at any age but are rare. The age at presentation varies and was stated as 19 years in one case. They can be in the form of cysts, sinus or fistula; cysts being the commonest. Common clinical presentation are pre-auricular swelling (24%), parotid swelling (36%) or cervical region swelling (41%). In our case, it was a fistulous opening that presented as discharging tract in upper part of neck. The Work classification system is widely quoted which classifies first branchial cleft anomalies into two types. Type 1 anomalies present as a cystic mass with ectodermal origin, with squamous epithelium but no skin adnexa or cartilage remnants. Type 2 present as cysts, sinuses, or fistula tracts and are of mixed ectodermal and mesodermal origin. They at times pass through the parotid either superficial or deep to the facial nerve, sometimes the tract may end in cartilage segment of external auditory canal. In our case, it was found to be traversing medial to facial nerve reaching up to the right external auditory canal. Imaging studies can help in determining the appropriate surgical approach. Prior drainage or rupture of cystic variant may lead to scar formation or fistula formation. The reported recurrence rate is as high as 14-22%. Definitive treatment for such anomalies is surgical excision where surgical plan needs to be tailored according to the tract traversed by the sinus or fistula. Key to complete excision is keeping the tract, cyst or fistula intact. Preservation of facial nerve is another key fragment in these surgeries for which sometimes the removal of superficial lobe of parotid gland is required with identification of facial nerve which is to be tailored specifically for each case.

CONCLUSION

First branchial cleft anomalies are rare but are a potential differential diagnosis when a patient presents with swelling over upper lateral neck or around the external ear. Management include early diagnosis, control of infection and complete excision with facial nerve preservation. Surgical approach should be based on the clinical examination, imaging and clinical course; and there is a need to safely identify facial nerve in almost all cases.

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