Odontogenic Choriostoma in buccal mucosa associated with suspected Goldenhar Syndrome

Prashant M Battepati1, Priyanka Gupta2, Bhar Sharanesha Manjunatha3* and Bheema Setty Manasali4

1Department of Preventive Dental Sciences, University of Taif, KSA
2Consultant Pediatric dentist, India
3Department of Oral Biology, Faculty of Dentistry, University of Taif, KSA
4Department of Fixed Prosthodontics, Faculty of Dentistry, University of Taif, KSA

Abstract

Presence of teeth or tooth like structures at unusual sites of perioral structures like maxillary sinus, nasal cavity, ear, lip etc have been occasionally reported. But presence of fully developed dental structures with supporting skeletal tissue at non tooth bearing areas is extremely rare entity. In this report a similar case of presence of supplemental dental tissue along with supporting skeletal structure at buccal mucosa affecting facial symmetry which was associated with suspected Goldenhar Syndrome is presented. Report of such condition is extremely scanty in available literature and has been termed as Odontogenic Choriostoma by certain authors.

Background

Presence of tooth or tooth like structures at extragingival sites is not uncommon. Various extra gingival sites are reported for presence of such kind of structures like nasal cavity, ear, tongue as well as mandibular condylar and coronoid process. Presence of such structures may cause malocclusion, difficulty in breathing, incomplete articulation and speech alteration. Some of them are diagnosed early whereas some remain subclinical and diagnosed incidentally on radiograph. Confirmatory diagnosis for such structures is made by co relating clinical, radiographic and histological examination.

Presence of tooth like structures at unusual site is considered ectopic and their presence in soft tissue is seldom observed. The term “choriostoma” has been proposed for such kind of structures which are histologically normal for a part of body except at the site where it is located [1]. Primary classification of choriostoma has been proposed based on its clinical and histomorphologic features like salivary gland choriostoma, osseous and cartilaginous choriostoma, glial and gastric choriostoma etc [2]. Osseous and cartilageous types of choriostoma are most commonly seen in oral cavity [1]. For presentation of such lesions with dental component as well as its location on extra gingival site, term 'odontogenic choriostoma' seems to be appropriate. We present an unusual case of an 8 year old girl with presence of tooth like structures in right buccal mucosa with supporting bony extension from right maxillary posterior region associated with suspected case of Goldenhar syndrome, for which term odontogenic choriostoma is used.

Case presentation

An 8 year old girl reported to department of Pediatric dentistry, with chief complaint of presence of decayed teeth in right and left back region of lower jaw with no associated symptoms.

On extra oral examination, loss of facial symmetry was evident (Figure 1). There was presence of diffuse swelling extending from right corner of mouth to tragus of right ear. Superiorly, it was extending to infraorbital region of right eye and inferiorly till angle of the mandible on same side. There was no obliteration of nasolabial fold seen. Right corner of the mouth was appeared to be stretched out and with presence of faded groove running (resembling a surgical scar) from right corner of mouth, which was merging with the cheek after travelling a distance of 1 cm. On palpation, swelling was non-tender and firm in consistency. On further questioning, parents revealed that the swelling was significant as she had visited pediatrician and ENT specialist when she was 2 years old. Reports by pediatrician revealed presence of Goldenhar syndrome, for which term odontogenic choriostoma is used.

Presence of proximal decay with respect to mandibular right first primary molar and left first primary molars.

While examining intra oral soft tissues, a supplemental tooth like structure was revealed on right buccal mucosa distal to permanent 1^st molar which closely resembled the morphology of permanent maxillary molar. With respect to tooth like mass, almost two third of crown was exposed in oral cavity which was facing permanent first molar (Figure 2). On palpation, surrounding mucosa was apparently normal and there were no signs of inflammation or infection. Bidigital palpation confirmed that the structure was firmly adherent to the underlying tissues as it was firm and immovable.

On further questioning, parents revealed that the swelling was first noticed few years back and gradually increased to present size. There was no history of trauma. Patient’s medical history was significant as she had visited pediatrician and ENT specialist when she was 2 years old. Reports by pediatrician revealed presence of Goldenhar syndrome, for which term odontogenic choriostoma is used.

*Correspondence to: Manjunatha BS, Associate Professor, Department of Oral Biology, Chairman, Scientific Presentation Committee, Member, Examination Unit, Faculty of Dentistry, Taif University, Kingdom of Saudi Arabia, E-mail: drmanju26@hotmail.com

Received: April 09, 2019; Accepted: April 25, 2019; Published: April 29, 2019
multiple ear abnormalities and ear tags, macrostomia, right eye anomalies and hearing impairment followed by advice for x-ray for cervical spine, echocardiography and test for hearing assessment. Investigatory reports by the department of audiology from a hospital were accessed which revealed that on performing BERA test (Brainstem Evoked Response Audiometry) there was moderate hearing impairment with respect to left ear and right ear was normal. Whereas, reports of ECG and chest x-ray had shown no abnormalities. One of the medical reports of the child also showed a provisional diagnosis of Goldenhar syndrome at the age of 2 years.

In order to get a pan oral view and to rule out presence of any additional teeth, OPG (Orthopantomogram) was advised. The OPG revealed mixed dentition stage (erupted permanent central incisors, molars and primary canines and molars) and the developing tooth buds of permanent canines, premolars and second molars along with two radiopaque masses with radio density comparable with neighbouring teeth in maxillary right quadrant (Figure 3). In addition a lateral cephalogram (Figure 4) was also made. Nevertheless due to superimposition, detailed morphology could not be ascertained in both the radiographs. Hence 3 dimensional imaging using Digital volumetric tomography (DVT) was advised for the right posterior maxillary region.

DVT images revealed presence of two tooth-like structures with well developed internal morphology (pulp chamber and root canals) located distal and buccal to maxillary right 1st permanent molar. One among them was partially erupted whereas other was completely embedded into bony stalk which appeared to be originated from posterior most part of maxillary bone (Figure 5).

Based on the available evidence the lesion was diagnosed as Odontogenic choriostoma and surgical excision of the lesion was planned, however surgery could not be performed as parents of the child did preferred to wait and delay the surgical intervention.

Discussion

This report has presented a case with erupted supplemental tooth in right buccal mucosa with bony stalk. Although, presence of this was a diagnostic enigma because of its unusual location and close resemblance with normal morphology of tooth.

There are various terminologies used in literature for such kind of peculiar structures like irregular eruption, ectopic tooth, ectopic odontoma [3] and odontogenic choriostoma [4]. The term ectopia is used to designate a condition where a tooth or tooth-like structure is located or erupts in a place remote from the normal site [3]. The present case showed eruption of tooth-like structure from the right buccal mucosa; hence the term ectopic can be used.

Noroozi and Arora described a similar case of mature tooth like molariform type in buccal mucosa with a bony stalk for which they proposed the term ‘odontogenic choriostoma’. The term odontogenic choriostoma by definition is a mass formed by the faulty development of tissue of a type not normally found at that site with inherent growth potential. Choriostoma have a composition limited to only 1 or two germinal layers [4].
The most common types of intra oral choriostoma are either osseous (osteomas) or cartilaginous (chondromas) and occur in the tongue [1]. They present as firm smooth sessile or pedunculated nodules usually symptomless, with increased predilection for women. There are only few cases have been reported of choriostoma with dental origin.

In 1971 Kroll et al published 9 cases of ectopic bony lesions of the oral cavity and called them osseous choriostoma [5]. A case of osseous choriostoma was found in buccal mucosa of a 28 years old woman with Klippel Feil syndrome, but no association with a tooth was reported and lesion consisted of lamellar bone, osteocytes and hematopoietic tissue [6]. In 1977 Bakry reported a case of 9 year old girl with Klippel Feil syndrome. The lesion consisted of a geminate tooth surrounded by follicle. The observation of in situ tooth formation in areas outside the dental arches has reflexively led to the primary consideration of odontogenic tissues are not seen at these sites [8]. It should be noted, that, with the exception of a single case of a tooth in the upper lip[9] or Wharton's duct[10], the buccal mucosa is only a known site of surgery near ear region. Hence author chose to refer this case as “suspected case” of Goldenhar syndrome.

In present case though patient’s history is significant with probability of goldenhar syndrome but when patient reported to us there were no such clinical distinguishing features suggestive of Goldenhar syndrome except for a surgical scar on right corner of mouth probably due to surgical correction of Macrostomia and history of surgery near ear region. Hence author chose to refer this case as “suspected case” of Goldenhar syndrome.

The case presented in this report was not associated with any history of trauma or familial tendency. The exact etiology of this lesion remains unknown. To our knowledge this is a first case presentation of unilateral odontogenic choriostoma consisting of multiple teeth like structures with associated bony stalk/ extension in a child with suspected case of Goldenhar syndrome.

References
1. Gopalakrishnan R, Koutras IG, Schauer GM, Schnitker G (2009) Dental (odontogenic) choriostoma of the tongue. J Oral Maxillofac Surg 67: 1135-1138. [Crossref]
2. Chou LS, Hansen LS, Daniels TE (1991) Choriostomas of the oral cavity: a review. Oral Surg Oral Med Oral Pathol 72: 584-593. [Crossref]
3. Venigalla A, Guttikonda LK, Nelaharhi H, Babburi S, Pinisetti S, et al. (2015) Ectopic compound odontoma in the buccal mucosa: report of a rare case. Case Rep Dent 2015: 835171. [Crossref]
4. Ide F, Mishima K (2012) What is an odontogenic choriostoma? J Oral Maxillofac Surg 70: 3-4. [Crossref]
5. Noroozi AR, Arora E (2011) Odontogenic choriostoma: report of a case. J Oral Maxillofac Surg 69: 3006-3009. [Crossref]
6. Krolls SO, Jaconway JR, Alexander WN (1971) Ossous choriostomas (osteomas) of intraoral soft tissues. Oral Surg Oral Med Oral Pathol 32: 588-595. [Crossref]
7. Gaitán-Cepeda LA, Quezada-Rivera D, Ruiz-Rodríguez R (2003) Osseous choristoma in intraoral soft tissues. Case report. Med Oral 8: 220-223. [Crossref]
8. el sedfy Bakry N (1977) An ectopic odontome in the cheek. Oral Surg Oral Med Oral Pathol 43: 583-584. [Crossref]
9. Ide F, Kikuchi K, Miyazaki Y, Mishima K, Saito I, et al. (2010) Keratocyst of the buccal mucosa: is it odontogenic? Oral Surg Oral Med Oral Pathol Oral Radiol Endod 110: e42-e47. [Crossref]
10. Rion CL (1930) Tooth in upper lip: report of case. J Am Dent Assoc 17: 2295.
11. Gupta DS, Tandon PN, Sharma S, Jurel SK, Majumder K (2011) Intraglandular tooth: a rare case report of tooth in submandibular salivary gland duct. J Oral Maxillofac Surg 69: e365-307. [Crossref]

12. West CB Jr, Atkins JS Jr (1988) Choristomas of the intraoral soft tissues. Otolaryngol Head Neck Surg 99: 528-530. [Crossref]

13. Sooksam M, Philipsen HP (1986) The intra-oral soft tissue osteoma: report of two cases. J Dent Assoc Thai 36: 229-234. [Crossref]

14. Long DE, Koutnik AW (1991) Recurrent intraoral osseous choristoma. Report of a case. Oral Surg Oral Med Oral Pathol 72: 337-339. [Crossref]

15. Dali M, Chacko V, Rao A (2009) Goldenhar syndrome: A report of a rare case. Journal of Nepal Dental Association 10: 128-130.