A Rare Case Report of Non-Hereditary Ectodermal Dysplasia with Impacted Teeth in Maxilla and Mandible in 18 Year Old Male.

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ABSTRACT

Ectodermal dysplasia is characterised by hypotrichosis, hypohidrosis and hypodontia. This X linked Mendelian type involving one or more ectodermal structures is usually seen in males rather than females. Etiology leads to disorder in the morphological aspect of cutaneous and oral embryonic ectoderm. We report a case of 18 year old male presenting features of ectodermal dysplasia with multiple impacted teeth in maxilla and mandible.

Keywords: Hypohidrosis, Hypotrichosis, Hypodontia, Ectodermal dysplasia.

INTRODUCTION

Defects in the derivatives of embryonic ectoderm involving two or more tissues lead to ectodermal dysplasia. Thurnam introduced the term in 1848 which was later coined by Weech in 1929. This X-linked inheritance, gene locus Xq11-21.1, is mostly seen in males rather than in females.[3-5] The dentition is affected due to its occurrence in the first trimester of pregnancy or even before the sixth week of embryonic life. Remaining ectodermal structures might be involved after eight weeks. Occurrence hints at frequency of 7 cases per 10,000 births.[2] Christ-Siemens Touraine syndrome and Clouston syndrome accounts for syndromal involvement. Skin, hair, nails, eccrine glands, and teeth are commonly involved.[8, 11] Here, we are present a case of ectodermal dysplasia in 18 year old male with multiple impacted teeth in maxilla and mandible.

CASE REPORT

A 18 year old male patient attended the Department of Dentistry with a chief complaint of absence of majority of teeth. Patient was conscious, oriented and afebrile. Vitals stable. Mouth opening and lateral temporomandibular joint movements were satisfactory. Hair distribution was sparse. Tolerance to heat was unsatisfactory. Skin, palms and soles were dry and parched. Protuberant lips, frontal bossing and depressed nasal bridge were also observed in extraoral examination. Vertical dimension of face was also reduced [Figure 1, Figure 2, Figure 3].

Figure 1: Left lateral view.

Majority of the teeth were missing during intra oral examination. Radiological investigations such as
orthopantomograph revealed multiple impacted teeth in maxilla and mandible [Figure 4]. Family history revealed that this condition has been observed only in this individual since generations on both parents side.

DISCUSSION

First type of ectodermal dysplasia is associated with hypotrichosis, hypohidrosis and hypodontia termed as Christ-Turaine syndrome, a X-linked type. Secondarily, Cloustan syndrome involving hair, nails and teeth excluding sweat glands is mentioned. Other syndromes depicting hereditary ectodermal dysplasia includes Rapp-Hodgkin Syndrome, Strandberg- Ronchese’s Syndrome, Rosseli-Gulienetti Syndrome, etc. The characteristic features include fine, sparse, lustreless fair hair in scalp, scaling of the skin, pyrexia, heat intolerance caused due to anomalies of the skin appendages, which include the hair follicles, sweat glands and sebaceous glands.\[6,7,9\] Frontal bossing, depressed malar prominence and nasal bridge, small size of the face, thick everted protuberant lips, wrinkled hyperpigmented, periorbital skin and large low set ears.\[10\] In our case, facial hyperpigmentation, frontal bossing, protuberant lips, decreased facial vertical dimension, normally set ears bilaterally, depressed nasal bridge followed by malar prominence, sparse hair, absence of alveolar processes, missing teeth and impacted teeth (seen radiographically) were presented. Differential diagnosis includes progeria, incontinentia pigmenti, congenital syphilis, idiopathic hypoparathyroidism – Addison’s disease – moniliasis syndrome, Rothmund-Thomson syndrome and Chondroectodermal dysplasia.

This condition attracts multidisciplinary approach of a dental surgeon, faciomaxillary surgeon, orthodontist and prosthodontist. Esthetics and function can be enhanced by prosthodontic measures such as implants.\[1\] Mainly, the absence of teeth might have a psychological impact on the individual. All these can be managed in a comprehensive manner if the condition is diagnosed early.

CONCLUSION

This condition is usually of hereditary nature, whereas in our case the family history played a major role in depicting that non-hereditary type exists as well.
REFERENCES


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