Prosthodontic Management of a Pediatric Patient with Christ-Siemens-Touraine Syndrome: A Case Report

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ABSTRACT

Ectodermal dysplasias (ED) are a group of rare genetic disorders characterized by congenital defects involving two or more ectodermal structures. Christ-Siemens-Touraine syndrome or hypohidrotic ectodermal dysplasia is the commonest form of ED. Hypohidrotic ectodermal dysplasia (HED) is an X-linked disorder characterized by excessively dry skin due to the absence of sweat glands (anhidrosis), sparse body hair especially on the scalp and eyebrows (hypotrichosis), brittle nails, absence of sebaceous glands (asteatosis) and malformed or absent teeth. Oral manifestations include oligodontia or complete anodontia, conical teeth, underdeveloped alveolar ridges, generalized spacing and delayed eruption of permanent teeth. This case report discusses a classical case of HED and the options for rehabilitation in a growing patient. A thorough knowledge about the clinical manifestations of ED will lead to proper diagnosis and appropriate treatment plan thereby leading to significant improvements in esthetics, phonetics and masticatory function in ED patients, which in turn leads to improved quality of life in these individuals.

Keywords: Christ-Siemens-Touraine Syndrome, Ectodermal dysplasia, Prosthodontic management.

INTRODUCTION

Ectodermal dysplasias (EDs) are a group of rare genetic disorders characterized by congenital defects involving two or more ectodermal structures.1 It was first described by Thurnam in 1848 and has an estimated incidence of 1 in 1,00,000 births.2,3 In 1921, EDs were linked to X chromosomes by Thadani.3

Ectodermal dysplasias are mainly of two types based on the function of sweat glands and its number namely Clouston syndrome or hidrotic ED and Christ-Siemens-Touraine syndrome or hypohidrotic ectodermal dysplasia (HED).4 Among the two types, the HED is the commonest. HED is an X-linked disorder characterized by excessively dry skin due to the absence of sweat glands (anhidrosis), sparse body hair especially on the scalp and eyebrows (hypotrichosis), brittle nails, absence of sebaceous glands (asteatosis), and malformed or absent teeth.5,6

Along with these features, the affected individuals might show other manifestations like heat intolerance, prominent supraorbital ridges, protuberant lips, and depressed nasal bridge. Oral manifestations include oligodontia or complete anodontia, conical teeth, underdeveloped alveolar ridges, generalized spacing, and delayed eruption of permanent teeth.6,7

In pediatric patients with HED, restoration of oral esthetics is vital, if not it can affect their psychological well-being. A complete oral rehabilitation of these patients is of paramount importance from a functional, physiologic, and psychosocial point of view.8,9 There are various treatment modalities for managing oligodontia or anodontia in pediatric HED patients, but the most frequently preferred is removable prosthodontics.10 Since these patients have underdeveloped alveolar ridges due to the lack of teeth, construction of dentures is more challenging than usual. This report describes an oral rehabilitation of a pediatric patient with Christ-Siemens-Touraine syndrome.

CASE DESCRIPTION

A 7-year-old boy presented with the chief complaint of missing teeth, difficulty in mastication, difficulty in speech, and lack of esthetics. On general examination, he had scanty eyebrows and eyelashes, dry and wrinkled skin, depressed nasal bridge, prominent supraorbital ridge, everted dry lips, and hyperpigmentation around the eyes (Fig. 1). Hair over the scalp were sparsely distributed (Fig. 2). Finger nails were brittle (Fig. 3). On oral examination, there was delayed eruption of permanent dentition, conical maxillary anterior teeth, and partial anodontia (Fig. 4). Bone ridge on the mandibular anterior region was very thin (Fig. 5). Panoramic radiograph showed evidence of tooth formation in both the jaws, and six tooth buds were present probably related to maxillary incisors and second permanent molars. All the second primary molars and first permanent molars were present. Four cone-shaped crowns suggestive of primary canines were also seen (Fig. 6).

During the physical examination, the patient showed no signs of mental problems and his vital signs were normal. His father said that his child was not able to sweat. He also stated that there was no production of tears when his child cried. Nobody in the

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patient’s family had similar condition and he was born of non-consanguineous marriage.

A sweat pore count test was done in which starch-iodine powder was applied to the dorsal skin. In this patient due to the lack of sweat, starch-iodine powder did not change color to deep purple and sweat pores were poorly visible.

Based on the case history, clinical examination, and investigations, a diagnosis of Christ-Siemens-Touraine syndrome
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or HED was made. Removable partial dentures were considered as the treatment option for this patient. Prior to impression making, composite buildup of the conical maxillary anterior teeth was done using strip crowns (Fig. 7). Primary impressions were made with alginate, and then custom trays were fabricated for functional impression. Acrylic bases with occlusal wax rims were made on the master casts for establishing maxillomandibular relations. Primary tooth forms were selected for better esthetics, maintenance of physiologic spaces, and to provide static occlusion. After the final insertion (Figs 8 and 9), routine oral hygiene instructions for the dentures like maintaining a soft diet for few days and to remove the dentures at night for promoting the healing of the oral soft tissue, etc., were given to both the child and his father. Periodic follow-up was advised for modification or replacement of the dentures taking into consideration the patient’s developing maxillary and mandibular ridge.

DISCUSSION

Christ-Siemens-Touraine syndrome or HED is the most frequently seen type of ED. It is said to occur due to any genetic defect in ectodysplasin signal transduction pathways. During morphogenesis, epithelial cells in developing teeth, hair follicles, nails, and sweat glands make use of this pathway and any genetic defect can lead to dysplasia, hypoplasia, or aplasia of these structures. Oral manifestations of these disturbances include hypodontia, tooth malformations, and hypoplasia of teeth. In the present case, the patient showed typical features of Christ-Siemens-Touraine syndrome in his face and oral cavity.

Prosthodontic rehabilitation at an early age in such cases has various advantages. It helps the child to adjust with the prosthesis and develop normal appearance. The phonetics, mastication, and the temporomandibular joint function are also developed. Removable partial dentures were considered as the best treatment option for this patient considering the patient’s age, need for mastication, phonetics, esthetics, and overall psychological well-being. During growth spurts, the prosthesis can be modified based on the developing alveolar ridges. Apart from oral benefits, an early prosthodontic intervention also imparts psychological benefits. But sometimes a “knife-edge” morphology as seen with the mandibular ridge in this case along with decreased alveolar bone height make oral rehabilitation an arduous task in such patients.

Prosthodontic rehabilitation of a patient with HED can include fixed, removable, or implant supported prosthesis. Fixed partial dentures are not commonly used in the treatment of HED because of the limited number of teeth (hypodontia) to act as abutments. There also a possibility of fixed prosthesis interfering with jaw growth in growing patients due to rigid connectors if the prosthesis crosses the midline.

Oral rehabilitation by dental implants is usually limited to patients who have completed craniofacial growth and is best advised to delay until adolescence. Implants placed in growing patients can cause several problems like the implant prosthesis being submerged and acting like an ankylosed tooth, restrictions on skeletal craniofacial growth and trauma to tooth buds. According to Yap and Klineberg, implants placed in ED patients before the age of 18 years have a higher risk of failure.

In the present case, dental implant therapy was not an option due to incomplete craniofacial growth and lack of alveolar bone support.

Therefore, in child patients and adolescents with HED, removable partial dentures provide the most logical reversible treatment option that enhances esthetics and function without threatening an already compromised dentition.
multidisciplinary treatment plan involving pediatric dentist, orthodontist, prosthodontist, and an oral-maxillofacial surgeon is absolutely vital for complete restoration of oral function. In the present case, since the patient is a growing child, periodical reviewing for the modification of the transitional removable appliance is required to compensate for the continued growth and development. Once the skeletal growth is complete, the patient can opt for a definitive prosthesis like implant supported dentures.

Dental practitioners are often among the first to identify these patients due to characteristic oral and extraoral features. Therefore, a thorough knowledge about the clinical manifestations of this condition will lead to proper diagnosis and appropriate treatment plan, thereby leading to significant improvements in esthetics, phonetics, and masticatory function in ED patients, which in turn leads to improved quality of life in these individuals.

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