Oral findings in West syndrome – A Case Report

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ABSTRACT

West syndrome is a severe form of epilepsy syndrome which is characterized by a triad of infantile spasms, characteristic EEG findings (Hypsarrhythmia) and developmental delay. Minimal literature is available on dental findings of West syndrome. This case report presents an eight year old male child with cryptogenic form of West syndrome having a history of multiple clusters of infantile spasms. Orodental manifestations of west syndrome have been described and its dental management has been discussed in this report.

INTRODUCTION

West syndrome, a severe form of epilepsy syndrome, is a triad of infantile spasms (typical age when seizures start), hypsarrhythmia and mental retardation [1]. The first detailed description of this condition was given by Dr. West in 1841 in his own four month old child as mentioned in the Lancet (1841) [2]. The condition is characterized by sudden flexion (bending forward) in a tonic (stiffening) fashion of the body, arms and legs. The incidence of West syndrome ranges from 2 to 3.5/10,000 live births [3,4]. Onset is generally in the first year of life. The peak age of onset is between 3 to 7 months, though onset at 4 years of age has been reported [5]. Spasms usually cease by 5 years of age, but other seizure types are reported in as many as 60% of children with West syndrome even after cessation of spasms [5]. West syndrome occurs in all ethnic groups, and is more common in boys than girls (ratio of 60:40) [6]. Although various papers have been published on West Syndrome from a medical standpoint, there are very few articles available in the literature on orodental findings and treatment. This article documents the case of an 8 year old male child diagnosed with West Syndrome with more emphasis on the orodental findings and their dental treatment.
CASE REPORT

An eight year old male child reported to the Department of Pediatric and Preventive Dentistry with the chief complaint of pain in the lower right back teeth region since 10 days. Detailed medical history of the child was taken. Patient was a known case of cryptogenic form of West syndrome and was diagnosed at the age of one and a half years. The child had normal motor developmental milestones till one year of age, then he had recurrent myogenic epilepsy and EEG findings showed hypsarrhythmia and infantile spasms which helped in his diagnosis. The last episode of convulsion was at the age of seven years three months. Patient had poor communication skills, delayed social skills and an aggressive behavior. Patient has been taking oral antiepileptic medication syp. Levetiracetam (Keppra) 2.5ml BD since last 7 years. The patient is the third child from a non-consanguineous marriage. Other two siblings were healthy and did not have any medical history. On physical examination, the neuromuscular delay was the most evident characteristic and there was atrophy of the upper and lower limbs (Figure 1 and 2). The child did not cooperate for thorough intraoral examination. Initially, symptomatic pain management was done with Syp.Paracetamol (125mg/5ml) and further dental treatment was planned under general anesthesia. The necessary blood and radiographic investigations were done and pediatric and neuromedicine opinions were taken. Written consent for dental treatment under general anesthesia was obtained from the parents and after evaluating all investigations, the patient was scheduled for dental treatment under general anesthesia.

The intra oral examination was done and intra oral periapical radiographs (IOPA) [Figure 7] were taken under general anesthesia. The intra oral examination showed that the patient had mixed dentition. 64,65,73,74,75,84,85 had carious lesions, out of which 74,75,84 and 85 had multisurface carious lesions [Figure 3 and 4]. On radiographic assessment, these teeth showed root resorption. Generalized attrition of all primary teeth was seen with hypoplastic white spots on the labial surfaces of 31 and 41 [Figure 5]. Angle's class I molar relationship was noted on both sides with deep arched palate and anterior open bite [Figure 6].

Glass ionomer cement restoration in 64,65,73 and extraction of 74,75,84 and 85 were done. Oral prophylaxis and topical fluoride varnish application was done [Figure 8 and 9]. The child recovered and accepted general anesthesia well.
Oral findings in West syndrome – A Case Report

Badnaware S et al.


Figure 4 - Preoperative view of mandibular arch

Figure 5 - White spot lesions on 31 and 41

Figure 6 - Intraoral view showing angle class I molar relationship

Figure 7 - IOPA – Upper row- mandibular anterior and maxillary anterior region and Lower row- mandibular left and right posterior region

Figure 8 - Postoperative view of Maxillary arch

Figure 9 - Postoperative view of Mandibular arch
DISCUSSION

According to etiologic classification, West syndrome is divided into two types i.e. Cryptogenic and Symptomatic [7]. The cryptogenic form of West Syndrome is mostly idiopathic, with an unknown cause; whereas the symptomatic variety can be attributed to prenatal, natal and post-natal causes [8].

Sparse information is available in the dental literature which emphasizes on the orodontal findings and dental management of West syndrome patients. Parents of children with special healthcare needs find it difficult to maintain good oral hygiene owing to the child’s low IQ level, limited communication skills and lack of manual dexterity [9]. Dentists need to place more emphasis on preventive dental care of such patients to avoid future dental problems. The importance of family commitment to daily oral hygiene and avoidance of a cariogenic diet should be encouraged.

Case reports on West syndrome documented by Regis[10], Neta[11] and Khatri et al. [12] showed similar intra oral findings i.e. multiple white spot lesions on labial surfaces, generalized tooth wear, altered chronology of teeth and gingival inflammation. Regis [10] found a deep arched palate, delayed eruption of permanent lower incisors and permanent first molars. Gingival enlargement is likely due to poor oral hygiene and white spot lesions are due to high sucrose diet and sugar containing medications. Neta[11] reported other oral findings like anterior open bite with tongue interposition in between arches, altered chronology, fissured tongue and low caries experience.

As there are many other conditions with similar orofacial findings like Rett’s syndrome, cerebral palsy, Angelman’s syndrome, Lennox gastaut syndrome etc., a dental surgeon should be aware regarding the clinical and oral presentation of West syndrome. Awareness about the importance of prompt diagnosis and timely treatment of patients with special health care needs should improve for better long-term outcomes. To provide treatment for patients with special health care needs, a dentist must evaluate each patient thoroughly in terms of personal characteristics, symptoms and behavior. The dentist should never make assumptions about the child’s degree of impairments without taking thorough medical and dental history. Parent or guardians should be interviewed for detailed history. Parental education and training is required to improve the oral hygiene of special children. This case report is of practical significance to the dentist in treating a patient with West syndrome.

CONCLUSION

As the oral health is a vital part of overall well being and general health, dental professionals should be included in the multidisciplinary health care team. Dentists should become familiar with the clinical features of West Syndrome in order to provide good dental care to patients.

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REFERENCES


