Case Report

Oromandibular-limb Hypogenesis Syndrome Type II C: A Rare Case

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Abstract

The oromandibular-limb hypogenesis syndrome comprises a group of anomalies which simultaneously affect the mandible, tongue, and maxilla with or without reductive limb anomalies. It is characterized by failure of development of the intraoral region and distal extremities. Multiple and variable deformities of the mandible, maxilla and tongue may occur in combination with a variety of limb defects. The wide range of presentation and combination of anomalies make classification difficult. They usually feature primarily in sporadic case reports because of their low incidence. The genetic origin of this syndrome is uncertain. It is congenital and there seems to be no sex predilection. The key radiographic features are retruded mandible, impacted teeth and malformed phalanges. When compared to available literature, frequently reported features like hypodontia, hypoglossia, microstomia, protruded maxilla and limb anomalies were present in our case. The case presented here is one of the rarest subtypes of this rare syndrome.

Key words: Hypoglossia, peromelia, teratogenic.

Introduction

Oromandibular-limb hypogenesis syndromes (OLHS) represent a group of rare conditions characterized by congenital malformations involving multiple sites such as the tongue, mandible, and limbs. In 1971, Hall classified OLHS into 5 major types and according to this, the case report presented here falls under type II C, which is hypoglossia-hypodactylomelia syndrome. The hypoglossia-hypodactyly syndrome, the Moebius syndrome, the Hanhart syndrome, the Charlie M syndrome and OMLH are possibly variants of the same condition, and it is often difficult to define the phenotypic boundaries between them. There is considerable overlap between these syndromes gathered under the term OLHS, with a marked variability of face and limb anomalies as well as other additional malformations. Limb deficiencies are major congenital malformations and can result from a number of etiological factors. Heat-induced vascular disruption has been considered as one of the etiological factors for these syndromes. Apart from this, teratogenic etiology has also been implicated. The genetic origin of these syndromes is uncertain. However, most of the cases are sporadic. Forty-seven cases of hypo-
glossia-hypodactylyia (Type I A) syndrome have been reported before 1990. The exact incidence of Type II C of OLHS could not be traced from the available literature. This case is reported to highlight the clinical and radiological presentation of this rare syndrome. The radiographic features consisted of retruded mandible in the lateral cephalograms, impacted teeth in orthopantomogram and malformed phalanges in wrist radiographs.

The most accepted classification was proposed by Hall in 1971.

Case report

A 23-year-old male patient reported to our institute with a chief complaint of speech difficulties since birth. The patient also complained of microstomia, malaligned teeth and difficulty in walking. The patient had no difficulty in breathing and swallowing. The patient was the second child out of two children. There was no history of consanguineous marriage between the parents. There was no history of similar findings in the family. The patient reported a history of drug intake by his mother during the third month of her pregnancy. On general examination, the patient had an unusual gait due to peromelia of lower limbs. The patient was well oriented to time and place.

On extraoral examination, the patient exhibited peromelia of upper and lower limbs (Figure 1a,b), microstomia, protruded upper anterior teeth, retruded chin, incompetent lips and fusion of the lower alveolar mucosa with the cutaneous part of the lower lip. Intraoral examination revealed constricted maxillary and mandibular arches, absence of lower labial sulcus due to the fusion of lower lip to the alveolar mucosa (Figure 1c), hypoglossia (Figure 1d), hypodontia and root stumps in relation to the teeth #36, #43 and #46. The orthopantomograph revealed thinning of both the condylar heads, asymmetry between the right and left mandibular bodies and increased gonial angle (Figure 2a). Lateral cephalogram revealed

Figure 1. Clinical photograph of the patient showing peromelia of upper limbs (a); hypodactylyia of lower limbs (b); constricted maxillary arch and fusion of the cutaneous part of the lower lip to the alveolar mucosa (c); clinical photograph showing hypoplastic tongue (d).
retruded mandible and protruded upper maxillary teeth (Figure 2b). The radiograph of the hand showed absence of carpal bones (Figure 2c). The radiograph of the leg showed hypoplastic tarsal bones skeletal pattern (Figure 2d). After a thorough evaluation by a team of medical specialists, we arrived at a final diagnosis of oromandibular-limb hypogenesis syndrome Type II C.

Discussion

Our patient presented with extraoral features, including convex profile, micrognathic mandible with relative maxillary prognathism and intraoral features, including microstomia, hypodontia, hypoglossia, and constricted maxillary and mandibular arches. Our patient also presented with upper and lower limb anomalies. A brief comparison of available literature with our case findings is discussed below.

Our patient presented with features like convex profile, retruded mandible and proclined maxillary anterior teeth, consistent with features reported by Wadhwani et al\textsuperscript{12} in a case report; however, they reported about bilaterally impacted teeth which were
not observed in our case. However, in rare instances protruded lower jaw has also been reported.\textsuperscript{13}

The case presented here also showed upper and lower limb anomalies similar to the findings reported by Wadhwan et al and Figuero et al.\textsuperscript{12,13}

It was first reported by Rosenthal\textsuperscript{14} in 1932 as aglosia congenita. The OLHS is a rare complex of jaw and limb defects with unclear aetiology.\textsuperscript{12} There seems to be no sex predilection.\textsuperscript{14} However, the proposed aetiology is heredity, maternal hyperthermia and positive drug history during pregnancy. This syndrome sometimes presents with cranial nerve palsy (sixth and seventh).\textsuperscript{6} Hermann et al\textsuperscript{15} analysed OLHS cases and found that there was severity of upper limb involvement, especially malformation of the feet, but they did not find cranial nerve palsies and this was significant in differentiating the cases. The group of patients with cranial nerve palsies included cases with limb defects similar to those observed in Hanhart syndrome and others with Poland anomaly; finally, cases with cranial nerve palsies without limb involvement were documented.\textsuperscript{15} No evidence of cranial nerve palsy was observed in the case presented here. Past medical history for exposure of the mother to drugs during pregnancy was positive in a case report.\textsuperscript{14} A similar history was reported in our case. Multiple site involvement and the wide range and combination of anomalies make classification difficult.\textsuperscript{16} There is overlap and similarity between different syndromic entities among similarities with OLHS, including a long list of syndromes like Moebius syndrome, hypoglossia hypodactyly syndrome, Hanhart syndrome, glossopalatine ankylosis syndrome, limb deficiency, splenogonadal fusion syndrome, and Charlie M syndrome. All are very uncommon except for Moebius syndrome.\textsuperscript{16} These groups of syndromes require a long-term and multidisciplinary approach.\textsuperscript{17} The case reported here is a rare syndrome with multiple site involvement. The treatment includes replacement of upper and lower limbs with prostheses, correction of malocclusion and speech therapy. Similar treatment modalities were suggested to the patient but the patient could not afford them.

\textbf{Conclusion}

The case presented here is a rare subtype of oromandibular-limb hypogenesis syndrome with oral manifestations, including hypodontia, hypoglossia, retruded chin and malaligned teeth along with limb anomalies. Almost all cases reported to date are seemingly sporadic.

\textbf{References}


\textit{Oromandibular-limb Hypogenesis Syndrome Type II C 139}

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