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HEMIFACIAL MICROSOMIA - A CASE REPORT WITH REVIEW OF **CASE REPORT LITERATURE** ^{1*}Sajad Ahmad Buch, ¹Subhas Babu, ¹Renita Lorina Castelino, ¹Shruthi Rao, ¹Kumuda Rao Hemifacial microsomia (HFM) is a congenital deformity characterised by unilateral deficiency in the volume of hard and soft tissue of face. It is a syndrome ABSTRACT predominantly affecting first branchial arch featuring poorly developed temporomandibular joint, mandibular ramus, muscles of mastication and ear. People with HFM may present with an underdeveloped ear (microtia) or absent ear (anotia) on the affected side of the face. The incidence of this disorder is 1:3000-26,000, which is usually seen at birth. Here we present a case of hemifacial microsomia in a 12-year-old girl with hearing difficulties on the affected side with review of literature. Key words; Hemifacial, microtia, syndrome Introduction **Case Report** ¹Department of Oral Medicine The occurrence of hemifacial microsomia has been A 12-year-old female patient reported to the and Radiology found to be about 1 in every 5600 live births and Department of Oral Medicine and Radiology with a A. B. Shetty Memorial thus one of the most significant craniofacial chief complaint of defect of the right side of the Institute of Dental Sciences disorders with accompanying facial asymmetry¹. face since birth. The patient was born full term to NITTE University, Hemifacial microsomia is the second most non-consanguineous parents. The patient had no Mangalore, India common defect of craniofacial structures after cleft significant family and antenatal history. The patient lip and palate. The term hemifacial microsomia complained of hearing deficit on the right side. was coined by a German physician Carl Ferdinand There was no family history of the similar von Arlt when he came across a case with an condition in other family members. There was no **Received:** asymmetrical face, eye, and ear in 1881. It was first history of prior intervention. The patient had 04.10.2016 reported as a condition in the 1960s with aural, oral consulted a dentist for a toothache a few months and mandibular developmental defects². It arises back and was advised analgesics for the same. On Accepted: due to developmental defects in first and second general examination, the patient was found to be 24.01.2017 branchial arches with unilateral underdeveloped well nourished with moderate built and normal bones of the jaws and associated soft tissues³. The mental status. On extra oral examination, there was deformities accompanying hemifacial facial facial asymmetry detected on the right side of the microsomia result due to destruction or disturbance face due to the defect since birth (Figure 1a). There of the neural crest cells⁴. According to Converse et was an absence of prominence over the right ramus al if cranial defects are associated with hemifacial and mandibular angle area due to the hypoplastic microsomia the condition should be called as right mandibular body. The mandible was retrognathic and deviated to right side upon craniofacial microsomia. **Corresponding author** opening (Figure 1b). The patient had a normal Sajad Ahmad Buch A variant of hemifacial microsomia having mouth opening with no clicking noticed in temporomandibular joints. The examination of additional features of vertebral anomalies and epibulbar dermoids in addition to unilateral muscles of mastication revealed no significant microtia, microsomia, mandibular condyle and findings. The right side of face showed a primitive ¹Department of Oral Medicine ramus malformation was described as Goldenhar and Radiology ear, microtia (Figure 1c) and there was a small syndrome by Gorlin et al^{2,4}. A. B. Shetty Memorial preauricular tag on left side of the face (Figure 1d). Institute of Dental Sciences The right side of the patient exhibited hearing NITTE University. A careful assessment of wide spectrum of deficit. Intraorally a V-shaped palate with high arch Mangalore, India abnormalities, etiology, and features associated was appreciated. There was crowding of teeth in with hemifacial microsomia is needed to manage the mandibular arch. The gingiva was soft and the functional incapacities and to improve oedematous with the maxillary right first aesthetics of patients with this disorder. We present permanent molar, and mandibular first permanent e- mail: a case of hemifacial microsomia with its buchh.sajad@gmail.com molars were grossly decayed. The patient exhibited characteristic clinical and radiographic findings Angle's Class I malocclusion bilaterally. Based on with an effort towards improving the knowledge the chief complaint and the clinical features a and an attempt to differentiate it from other similar provisional diagnosis of hemifacial microsomia of syndromes. right side was made. Hypoplasia of the ramus,

condyle, coronoid process and mandibular body on the right side was evident on panoramic radiograph along with prominent antegonial notch. The absence of third molar tooth buds in first, third and fourth quadrants could also be appreciated on panoramic radiograph (Figure 2a).



Figure 1. Clinical pictures:

Figure 1a. Clinical photograph of the patient showing defect on the right side of face.

Figure 1b. Clinical photograph of the patient showing deviation of mandible on the right side on opening.

Figure 1c. Clinical photograph of the patient showing microtia of the right ear.

Figure 1d. Clinical photograph of the patient showing ear tag on the left side.

The posterior anterior view of the skull showed a reduced height of ramus on the right side along with a shift of midline towards the right side. The nasal septum and maxilla were also seen to deviate towards the right side (Figure 2b). Lateral cephalogram showed an underdeveloped mandible along with a steep mandibular plane with a marked reduction in the height of ramus (Figure 2c). The patient was referred to an otolaryngologist for the hearing deficit. The patient was advised extraction of grossly decayed teeth and oral prophylaxis. The patient was also advised corrective surgery and correction of malocclusion, ear prosthesis for ear abnormalities and use of hearing aids to facilitate hearing.



Figure 2. Radiographic images:

Figure 2a. Panoramic radiograph showing hypoplasia of the ramus, condyle, coronoid process and mandibular body on the right side.

Figure 2b. Posterior-anterior view of skull showing reduced height of ramus on the right side along with a shift of midline towards the right side.

Figure 2c. Lateral cephalogram showing an underdeveloped mandible along with a steep mandibular plane.

Discussion

Hemifacial microsomia (HFM) is a congenital disorder of rare entity with wide phenotypic diversity. HFM has been seen to affect males more than females usually in the ratio of 3:2 and right side of the face is more affected than left side⁵. The case presented here is of a female patient with right side involvement. Early loss of neural crest has been reported to result in certain factors leading to the clinical presentation of hemifacial microsomia⁴. Certain teratogens, gene defects and anomalies of the vascular system, acting individually or collectively cause derangement of usual normal development and in turn result in features seen in these patients. Poswillo postulated that maternal intake of 10mg/kg Thalidomide causes rupture of the stapedial artery and results in its total or incomplete development. Derivatives of first and second branchial arches are also seen to suffer localized necrosis. According to researchers, this could lead to compromised blood supply to first and second branchial arches during the first 6-8 weeks of pregnancy^{4,6}. The characteristic diagnostic features of hemifacial microsomia include asymmetrical hypoplasia of facial structures like hypoplasia of mandible, microtia, and presence of preauricular tags with or without periauricular skin tags⁷. The shifting of the chin and facial midline towards the affected side results in an oblique lip line with the corner of the mouth higher on one side than the other⁸. According to some authors, it is mandatory to have unilateral microtia or any kind of ear abnormality such as preauricular tags in such cases to be diagnosed as hemifacial microsomia⁴. Our case presented with the features of unilateral mandibular hypoplasia because of reduction in height of mandibular ramus, a shift in the midline resulting in oblique lip line with one corner of the mouth placed higher than the other and unilateral microtia which helped us to arrive at the diagnosis of hemifacial microsomia. The present case also had a preauricular tag on the normal side of the face. Treacher-Collins syndrome a different syndrome shows bilateral ear abnormalities in addition to micrognathia⁹. This could have led to a diagnostic confusion

in our case but as there was neither a sign of hypoplasia of zygomatic arches nor any downward slanting of palpebral fissures, therefore we diagnosed our case as hemifacial microsomia. Such distinction helps to supplement any efforts and improves knowledge about the wide spectrum of features associated with hemifacial microsomia. The ear canal in HFM patients is either absent or its end is blinded on the affected side¹⁰. The middle ear abnormalities result in conduction defects in 30 to 50% of HFM cases ^{3,4}. This feature coincided with our case. Various features on intraoral exam seen in HFM cases include missing teeth on affected side (mandibular second premolar), absent mandibular third molars, delay in tooth development and susceptibility for aplastic and hypoplastic teeth^{10,11}. The present case showed the absence of third molar tooth buds in first, third and fourth quadrants. The various advanced imaging tools used for diagnostic and treatment purposes in HFM include Cone Beam Computed Tomography (CBCT), Multi- Slice Computed Tomography (MSCT), Magnetic Resonance Imaging (MRI) and Three Dimensional (3D) Surgical stimulation models⁸. The aim of surgery is to correct facial asymmetry and restore function¹². There is a need for collaborative efforts between different medical specialities for the wide spectrum of disabilities involving the various structures of the craniofacial area and occasionally other regions of the body such as associated defects in kidneys or heart in the management of HFM¹³. The management should begin early in life and should involve a team of specialists which include geneticists, audiologists, speech pathologists, physicians, plastic maxillofacial surgeons, surgeons, pedodontists, prosthodontists and orthodontists⁵. The management of HFM involves synergistic approach from specialists of different medical disciplines. Pre-and post-growth phase surgeries, use of grafts for deficit regions of the craniofacial framework, osteotomy for protraction of deficient mandible in mild cases of micrognathia of the mandible are among the treatment options in the management of HFM. Costochondral grafts can be given to provide new growth sites and soft tissue augmentation of the involved side can be done using the procedures of genioplasty and microvascular free flaps¹⁴

Conclusion

Hemifacial microsomia is a developmental abnormality in which there is a deficiency in both soft and skeletal tissues of the maxillofacial region on one side of the face. Early diagnosis and treatment should be planned with a team of experts which include dental radiologists, orthodontists, pedodontists, speech therapist, surgeons and paediatric surgeons for efficient clinical care to restore normal functions in the patients affected with hemifacial microsomia.

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