Crouzon’s Syndrome: A Rare Genetic Disorder

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ABSTRACT
Crouzon’s syndrome, also known as brachial arch syndrome, is an autosomal dominant disorder with complete penetrance and variable expressivity. Described by a French neurosurgeon in 1912, it is a rare genetic disorder. Crouzon’s syndrome is caused by mutation in the fibroblast growth factor receptor 2 (FGFR2) gene. Normally, the sutures in the human skull fuse after the complete growth of the brain, but if any of these sutures close early, then it may interfere with the growth of the brain. The disease is characterized by craniosynostosis, with associated dentofacial anomalies. This report describes the different clinical features in a 10-year-old male patient, with particular reference to characteristic findings of this syndrome.

Keywords: Crouzon’s syndrome, Fibroblast growth factor, Premature synostosis.

INTRODUCTION
Cranial skeletogenesis is a unique cranial malformation, although uncommon, compromising not only the function but also the mental well-being of the person.1 Described by a French neurosurgeon, Octave Crouzon, in 1912,2 it is a rare genetic disorder. It may be transmitted as an autosomal dominant genetic condition. Crouzon’s syndrome is caused by mutation in the fibroblast growth factor receptor 2 (FGFR2) genes.3

The disease is characterized by premature synostosis of coronal and sagittal sutures, which begins in the first year of life.4 Features of the condition include craniosynostosis, midfacial hypoplasia, maxillary retrusion, hypertelorism, and shallow orbits, which give rise to ocular proptosis.5 Intraoral manifestations include mandibular prognathism, overcrowding of upper and lower teeth, and V-shaped maxillary dental arch.2 Narrow, high, or cleft palate and bifid uvula can also be seen.

Occasional oligodontia, macrodontia, peg-shaped, and widely spaced teeth have been reported.4
Crouzon’s syndrome occurs in approximately 1 in 25,000 births worldwide.6 Crouzon’s syndrome makes up approximately 4.8% of all cases of craniosynostoses.7 No known race or sex predilection exists.4 The differential diagnosis of Crouzon’s syndrome includes simple craniosynostosis as well as Apert syndrome, Carpenter syndrome, Saethre–Chotzen syndrome, and Pfeiffer syndrome.8 The various abnormalities associated with Crouzon’s syndrome have been listed in Table 1.9

Table 1: Abnormalities associated with Crouzon’s syndrome

| Cranium          | Brachycephaly and acrocephaly |
|------------------|--------------------------------|
| Craniosynostosis | Palpable ridge                  |
| Flat occiput     | Frontal bossing                 |
| Frontal bossing  | Facial features                 |
| Maxillary retrusion |                          |
| Malar deficiency | Relative mandibular prognathism |
| Ear              | Low set ear                     |
| Conductive hearing loss |                          |
| Bilateral atresia of auditory meatus |                  |
| Eye              | Downslanting palpebral fissure  |
| Hypertelorism    | Exophthalmos                    |
| Hypertelorism    | Iris – coloboma                 |
| Hyperelorism     | Ptosis                          |
| Nystagmus        | Exposure keratitis              |
| Nose             | Beaked nose                     |
| Beaked nose      | Deviated nasal septum           |
| Mouth            | Short upper lip                 |
| High arched and narrow palate |                          |
| Cleft palate and bifid uvula |                      |
| Neurological     | Meningeal disease               |
| Headache         | Meningeal disease               |
| Mild to moderate mental retardation |                  |
| Seizures         | Cervical spine abnormalities (scoliosis) |
| Respiratory      | Calcification of stylohyoid ligament |
| Breathing        | Ménière’s disease (vertigo, dizziness, and/or ringing in the ear) |
| difficulty       | Respiratory system              |
| Sleep apnea      | Breathing difficulty            |
| Cutaneous        | Tonsillar hypertrophy           |
| Acanthosis nigricans |                                |
CASE REPORT

A 10-year-old male patient, born by a full-term normal delivery, reported to our department along with his father. The chief complaint as reported by the father was pain in relation to the upper left tooth region of the child for the past 1 week. It was the patient’s first dental visit. Since the child’s appearance and head size were not normal, the family and medical history were taken in detail. No history of any systemic illness or drug allergy was reported by the patient’s father. There were no reported anomalies in any siblings or near relatives.

EXTRAORAL EXAMINATION

Clinical examination revealed hypertelorism, external strabismus, optical atrophy, hypoplastic malar process, hypoplastic maxilla, relative mandibular prognathism, depressed nasal bridge, deviated nasal septum, short upper lip, and long philtrum (Fig. 1). No digital abnormalities were present. Extraoral examination revealed elliptical-shaped head, with dolichofacial growth pattern, and concave facial profile (Fig. 2).

INTRAORAL EXAMINATION

Intraoral examination revealed that the patient was in the late mixed dentition phase. There was apparent class III malocclusion with edge-to-edge incisal relationship with posterior crossbite on the right side and edge to edge on the left side (Fig. 3). Patient had V-shaped maxillary arch (Fig. 4), with crowding present especially in the lower labial segment (Fig. 5). The child also had high arched palate and bifid uvula, which are the characteristics of this syndrome (Fig. 6). Upper right deciduous second molar (55) and upper left deciduous canine (63) were missing. The tooth in question, i.e., upper left deciduous first molar (64), was found to be carious and grade II mobile (Fig. 4). Oral hygiene status was poor. Grade III calculus was evident, especially in the lower labial segment (Fig. 5).

TREATMENT PLAN

First, intraoral periapical radiograph with respect to 64, lateral cephalogram, and lateral skull and posteroanterior skull views of the patient were advised; 64 region showed physiologic resorption of the concerned tooth (Fig. 7),

Fig. 1: PA Skull view and Lateral skull view showing hypoplastic maxilla, relative mandibular prognathism and cerebral impressions

Fig. 2: Front profile photograph showing depressed nasal bridge and deviated nasal septum

Fig. 3: Right lateral profile photograph showing concave profile and relative mandibular prognathism

Fig. 4: Intraoral photograph showing edge to edge incisal relationship
and extraction of 64 was advised. Radiovisioigraphy of 55 tooth region (Fig. 8) did not show bone overlying the erupting premolar, hence there was no need for the space maintainer in that region. Radiographic findings revealed hypoplastic maxilla, relative mandibular prognathism (Fig. 9), cerebriform impressions, hypophyseal fossa enlargement, and cervical region abnormalities, i.e., fusion of posterior bodies and elements (C2–C3).
(Fig. 10). After clinical and radiographic investigations, complete oral prophylaxis was done, and the patient was advised to rinse twice daily with chlorhexidine mouthwash. Then extraction of 64 was performed under local anesthesia, and postextraction instructions were given with particular emphasis on lip biting, which is very common among children after being anesthetized. Also, ear, nose, and throat and ophthalmology opinion for the patient was taken. Patient was diagnosed to have congenital dacryocystitis along with slight visual impairment. Figures 11 and 12 show follow-up photographs. Digital panoramic radiograph of the patient was done at 6 months recall, which revealed normal eruption of the succedaneous teeth.

**DISCUSSION**

The phenotypic features of Crouzon’s syndrome may be absent at birth and may evolve gradually during the first few years of life. It is commonly inherited as an autosomal dominant trait, with complete penetrance and a variable expressivity, but about one third of the cases do arise spontaneously. The male to female preponderance is 3:1. With the advent of molecular technologies, the gene for Crouzon’s syndrome could be localized to the FGFR2 gene, at the chromosomal locus 10q25.3-q26, and more than 30 different mutations within the gene have been documented in separate families. Hypertelorism was a universal finding in the affected individuals and is thought to arise due to a decrease in growth of the sphenozygomatic and sphenotemporal sutures. The appearance of an infant with Crouzon’s syndrome can vary in severity from a mild presentation with subtle midface deficiency to severe forms with multiple cranial sutures fused and marked midface and eye problems. Upper airway obstruction can lead to acute respiratory distress and the presence of mental retardation is rare in these children.

**CONCLUSION**

An understanding of these abnormalities is necessary for the dental team to make the appropriate referrals to ensure that the patient receives the best available care. The pediatric dentist should be an integral part of the multidisciplinary team.

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