Apert’s Syndrome- A Rare Craniofacial Anomaly

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Abstract

Introduction: Apert’s Syndrome is a rare autosomal dominant disorder characterized by craniosynostosis, craniofacial anomalies, and severe symmetrical syndactyly (cutaneous and bony fusion) of the hands and feet. Other craniosynostosis syndromes such as Carpenter syndrome, Crouzon disease (craniofacial dysostosis), Pfeiffer syndrome and Saethre-Chotzen syndrome have clinical features that are similar to Apert’s syndrome. Case Presentation: The case of a 10 month old infant with Apert’s syndrome is presented here. The characteristic features of Apert’s Syndrome are discussed. Conclusion: A multidisciplinary team approach comprising of neurosurgeons, craniofacial surgeons, plastic surgeons, otolaryngologists, orthodontists, orthopaedic surgeons ophthalmologists, radiologists, geneticists, paediatricians, clinical psychologists, speech and language pathologists is needed for the effective management of this condition.

Key words: Craniofacial anomaly, premature synostosis, syndactyly

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Introduction

Apert’s Syndrome (Acrocephalosyndactyly Type I), is a rare congenital disorder characterized by premature fusion of cranial sutures (craniosynostosis), malformations of skull, hands, face and feet [¹, ²]. It is a branchial arch syndrome affecting the first branchial arch, the precursor of maxilla and mandible. Apert’s syndrome is caused by mutations in a gene called fibroblast growth factor receptor 2 which is located on chromosome 10 [³, ⁴]. The mode of inheritance is reported to be autosomal dominant and associated with advanced paternal age. However, sporadic occurrence of many cases has been reported, possibly representing new mutations [⁵]. This rare clinical abnormality has to be differentiated from other craniofacial syndromes such as
Carpenter syndrome (Kleeblattschadel, cloverleaf skull deformity), Crouzon disease (craniofacial dysostosis), Pfeiffer syndrome and Saethre-Chotzen syndrome. The case of a 10 month old infant with Apert’s syndrome is reported here. The characteristic clinical features as well as multidisciplinary approach in the management of the case are discussed.

**Case Report**

A 10 month old infant presented with increased distension of head and fused digits of the hands and feet. The infant was a premature baby born at 8 months of gestation.

The baby developed impaired vision, abnormal shape of the skull, with mild exophthalmos of eye, and delayed milestones. The child failed to recognize objects. There was a delay in gross and fine motor skills as well as speech and language development. On examination there was facial disfigurement with frontal bossing.
Figure 1: Clinical Photograph of the infant showing high arched palate, cleft of soft palate extending till the hard palate and syndactyly.

Plain X-ray of the skull anteroposterior (AP) view showed obliteration of coronal and lambdoid suture lines with obvious bony continuity, mild ocular hypertelorism and crowding of teeth. Plain X-ray of the hand AP view showed the fusion of the phalanges. Plain X-ray of the feet revealed the bony fusion of first and second metatarsals (Figure 2).
Figure 2: Plain X-ray of hands and feet showing fusion of phalanges and metatarsals
Computerized tomography (CT) scan of the skull with 3–dimensional reconstruction showed increase in biparietal diameter with fusion of coronal sutures and partial fusion of lambdoid sutures suggestive of craniosynostosis causing brachycephaly. Partial agenesis of corpus callosum was also seen. Anterior and Posterior fontanella were widely separated (Figure 3, 4, 5). Mild prominence of frontal horn of lateral ventricle was also seen.

Figure 3: CT scan of the skull with 3–dimensional reconstruction showing flat occiput with fusion of coronal sutures and partial fusion of lambdoid sutures
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Figure 4: CT scan of the skull with 3–dimensional reconstruction showing craniosynostosis with wide anterior fontanelle

Figure 5: CT scan of brain showing partial agenesis of corpus callosum
Surgical management has been planned in several stages in order to (i) prevent further closure of coronal sutures and damage to the developing brain (ii) facilitate proper speech and (iii) to provide aesthetically and functionally acceptable life.

In this case, fronto-orbital advancement was done along with the release of the coronal and lambdoid sutures to permit normal growth of the brain and ventriculoperitoneal shunt was done to reduce hydrocephalus in **Stage 1**. This was followed by two flap palatoplasty (Bardach’s technique) for management of cleft palate in **Stage 2**.

The following treatment is being further planned:

**Stage 3:** Le Fort III osteotomy and maxillary distraction advancement to correct the midfacial hypoplasia

**Stage 4:** Management of syndactyly by releasing the first and fourth web space early. This will be followed by lengthening of the thumb to increase the first web space. The second and third web spaces will be then released in order to establish a fully functioning hand. The aim is to make it possible for the child to grasp with his hands.

**Stage 5:** Correction of Ophthalmic and ear anomalies

The patient has improved with frontoorbital advancement and ventriculoperitoneal shunt. Patient is having proper growth of skull subsequent to surgery. The child has started to develop normal speech. Surgery for midfacial hypoplasia and syndactyly is being planned in the near future.

**Discussion**

Apert’s syndrome (Acrocephalosyndactyly Type I) is a rare congenital anomaly described by Eugène Charles Apert, a French pediatrician, in 1906, in nine people sharing similar attributes and characteristics [1]. It has a prevalence of 15.5 per million live births and accounts for 4.5% of all cases of craniosynostosis [7].

There is a considerable overlap between the features of Apert’s syndrome and other craniofacial syndromes particularly Carpenter syndrome (Kleeblattschadel, cloverleaf skull deformity), Crouzon disease (craniofacial dysostosis), Pfeiffer syndrome and Saethre-Chotzen syndrome [6].

In Apert’s syndrome, a unique fibroblast growth factor receptor 2 (FGFR2) gene mutations, that map to chromosome bands 10q25–q26 has been described. These mutations lead to an increase in the number
of precursor cells that enter the osteogenic pathway. Premature osteogenesis leads to fusion of sutures and the characteristic skeletal abnormalities. Once a suture becomes fused, growth perpendicular to that suture becomes restricted, and the fused bones act as a single bony structure. Compensatory growth occurs at the remaining open sutures to allow continued brain growth. However, complex, multiple sutural synostosis frequently extends to premature fusion of the sutures at the base of the skull, causing midfacial hypoplasia, shallow orbits, a foreshortened nasal dorsum, maxillary hypoplasia, and occasional upper airway obstruction [3,4,9,10].

The diagnosis of Apert’s syndrome is mainly dependent upon the clinical and the radiological findings since the molecular analysis for detection of the specific mutation is expensive. Apert’s syndrome is characterized by craniosynostosis, midfacial hypoplasia and syndactyly of hands and feet with a tendency for fusion of bony structures. Premature fusion of coronal sutures results in an acrocephalic (cone-shaped) head with flat occiput, short anterior-posterior diameter, prominent elongated forehead and a short broad nose with a bulbous tip and reduced nasolabial angle. Midfacial hypoplasia leads to retrusion of the middle third of the face and relative mandibular prognathism. Shortening of the bony orbit contributes to ocular features such as hypertelorism, proptosis, strabismus and down-slanting palpebral fissures [8]. The patient reported here had most of these features.

The oral manifestations of Apert’s syndrome include lateral swelling of the palatine processes, a cleft uvula, tooth crowding, bulging/thickened alveolar processes, and delayed dental eruption. The palatine process swelling may be due to accumulation of mucopolysaccharides which sometimes leads to pseudoclefting of the palate [11]. The prevalence of a real cleft palate is reported to be seen in 25-75% of Apert syndrome subjects [12]. In this case, the patient had submucous cleft in the soft palate which extended to the hard palate.

Skull radiography reveals craniosynostosis which usually involves coronal sutures [13]. In this case, apart from fusion of coronal sutures, partial closure of lambdoid sutures was also present at 10 months itself. Hand radiography shows characteristic finding of complete syndactyly involving the second and fifth digits (mitten hands). Radiography of the feet can be
performed to evaluate for cutaneous and osseous syndactyly. The characteristic finding is complete syndactyly involving the second and fifth digits (sock feet). The patient reported here had syndactyly of both hands and feet.

Computerised tomography (CT) scan with comparative 3-dimensional reconstruction analysis of the calvaria and cranial bases has become the most useful radiological examination in identifying skull shape and presence or absence of involved sutures. CT scan precisely reveals the pathological anatomy and permits specific operative planning. Magnetic Resonance Imaging is the imaging modality of choice for detecting intracranial abnormalities. The intracranial malformations reported on neuroimaging include ventricular enlargement, corpus callosum hypoplasia, septum pellucidum hypoplasia, cavum vergae and arachnoid cysts. Many patients develop normal intelligence although mental retardation has been reported in some. This has been attributed to intracranial malformations, increased intracranial pressure or family environment. Partial agenesis of corpus callosum with prominence of lateral horn of ventricle was noted in the CT scan in the case reported here.

Presence of cleft palate could predispose patients to recurrent ear infections besides causing speech difficulties. Psychosocial, speech and hearing assessment is important in patients with Apert’s syndrome to undertake rehabilitative measures to prevent cognitive impairment and delay in speech and language development.

The main modality in the management of Apert’s syndrome is surgery. The main aims of surgery in Apert’s syndrome are to:

- Release the cranial sutures in order to permit a proper brain development
- Prevent hydrocephalus due to raised intracranial tension
- Repair the cleft in order to prevent nasal regurgitation as well as facilitate proper speech
- Correct midface hypoplasia to facilitate proper breathing and improve aesthetic appearance
- Release of syndactyly to provide a better grasp.

The other modalities of treatment include application of tear drops to prevent corneal ulcer and correct the refractive errors. The overall aim is to give the patient a functional
and socially acceptable life. Although. The timing of surgery is controversial, early surgical treatment for cranial malformations before the first year of life is suggested to minimize cognitive loss and maximize normal development \[15,16\]. A strong multidisciplinary team comprising of neurosurgeons, craniofacial surgeons, plastic surgeons, otolaryngologists, orthodontists, orthopedic surgeons, ophthalmologists, radiologists, geneticists, paediatricians, clinical psychologists, speech and language pathologists is necessary for the care of these patients to improve their quality of life.

**Conclusion**

It is essential to be aware of the clinical manifestations of Apert’s syndrome so as to facilitate diagnosis and initiate appropriate treatment strategies in order to give the patient better quality of life. Effective management needs the collective efforts of a multidisciplinary team comprising of health care professionals. Counseling of parents regarding the necessity for treatment and motivating them to provide necessary support and family environment for the psychosocial development of these patients is also essential.

**References**

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