Case Report

Radial club hand managed with ulnar osteotomy and centralization of hand: a case report and review

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ABSTRACT

Radial club hand also called radial longitudinal deficiency or radial dysplasia is a preaxial longitudinal failure of formation. As the defect is preaxial it is often associated with thumb hypoplasia or anomaly of the radial aspect of the carpus. It is diagnosed clinically and on X-rays. It is frequently syndromic so it is a must to look for associated congenital anomalies by doing a thorough clinical examination. The frequency of this anomaly is between 1:50000 to 1:100000 live births. The incidence of all radial ray-deficient limbs, including hypoplastic thumbs alone, is approximately 1:30000. The radial deficiency is bilateral in 50% of the cases and the male:female is 3:2. It includes a wide spectrum of disorders that encompass an absent thumb or thumb hypoplasia, a thin first metacarpal and an absent radius. We report here a 1.5 years old child with isolated type IV radial club hand without any restricted range of motion in elbow managed with osteotomy of ulna and centralization of hand.

Keywords: Radial club hand, Ulnar osteotomy, Hand

INTRODUCTION

Club hand deformities can be either preaxial or postaxial. Preaxial is radial defect whereas postaxial is ular defect. Radial club hand or radial dysplasia is an uncommon congenital anomaly. It is a longitudinal deficiency along the preaxial or the radial aspect of the upper extremity. The frequency of this anomaly is between 1:50000 to 1:100000 live births. The incidence of all radial ray-deficient limbs, including hypoplastic thumbs alone, is approximately 1:30000. The radial deficiency is bilateral in 50% of the cases and the male:female is 3:2. It includes a wide spectrum of disorders that encompass an absent thumb or thumb hypoplasia, a thin first metacarpal and an absent radius. The radial ray which includes radius, scaphoid, trapezium, thumb and associated soft tissue structures of which all or part can be missing depending on severity. The median nerve lies very superficial at the level of wrist. Other muscles in upper limb may be absent, abnormal or accessory like presence of brachioarcapalis in forearm, consistently found in patients with TAR syndrome. The radial artery is absent in 86.5% of the cases with a normal ulnar artery. A significant of cases have a persistent embryogenic median artery of the upper limb. Several theories are postulated like maternal exposure to valproic acid, compression of the uterus and vascular injury, but the current theory relates the aetiology of the radial club hand to the apical ectodermal ridge (AER) which is a thickened layer of ectoderm that directs the differentiation of the underlying mesenchymal tissue and limb formation. Removal of a portion of AER in chick embryos has produced anomalies which are similar to the radial club hand. The extent of the deformity is related to the degree and extent of AER absence. Variuos gene defects have been identified in syndromic associations like mutation of TXB in chromosome 12 in Holt-Oram syndrome, two abnormal FANC gene in Fanconi anaemia. Heikel, based on the amount of the radius which was present, classified
radial dysplasia into four types, ranging from a present but
defective distal radial epiphysis (Type I) to a complete
absence of the radius (Type IV). TYPE I: mildest form
with defective distal radial epiphysis, TYPE II: Involves a
limited growth of the radius on both its distal and proximal
sides, TYPE III: Absence of two-thirds of the radius, most
commonly the distal side and TYPE IV: Complete absence
of the radius along with a complete or a near complete
absence of the thumb, which is the most common and most
severe longitudinal deficiency. 

Treatment is usually based on age with splinting being
effective and well tolerated till 6 months of age. Beyond
this till 18 months a stretching program for the parents to
the child is recommended combined with night splinting.
Stretching and splinting should be started early in infancy
to maintain soft tissue length. Beyond 18 months
conservative procedures are not effective and surgical
intervention is done. Operative procedure includes use of
a bilobed flap, soft tissue release, ulnar osteotomy, and
temporary longitudinal wiring of the carpus to the distal
end of the ulna in the corrected position as done in our
case. Hypoplastic thumb is managed either by web space
lengthening for Blauth type I defects, additional
opponensplasty and ulnar collateral ligament
reconstruction for type 2, tendon transfer for type 3A. For
Blauth 3B, 4 and 5 we wait till age of 2 years and perform
policization of a functional index finger. Other methods
include radialisation which in itself is a modification of
centralization with radial sided tendons transferred
ulnarily. Ilizarov correction can be used to lengthen ulna
through osteotomy in conjunction with realignment
procedures. Vascularised epiphysyal transfer of a
metatarsophalangeal joint unit from foot to the radial side
of wrist is done via microsurgical techniques. Surgically
acceptable postoperative period may not be satisfactory in
long run due to recurrence of deformity or arrest of distal
ulnar growth brought on by surgery. Contraindications for
the operative treatment includes life threatening anomalies
which are not compatible with long life, ulnohumeral
synostosis, mild deformity with adequate radial support
(type I and some type II), and older patients who have
accepted the deformities and have adjusted accordingly. 

So, keeping in mind the above facts, the present case report
and review was conducted to manage the radial club hand
with ulnar osteotomy.

CASE REPORT

A 1 year 6 months old male child was brought to the OPD
at Pt. B. D. Sharma PGIMS, Rohtak, Haryana, with a
history of a deformed left upper limb in the form of a short,
curved, left forearm and a hypoplastic thumb since birth.
He was a product of a non-consanguineous marriage and
his perinatal history was uneventful. There was no history
of blood transfusion. There was no family history of a
similar deformity in the past two generations. He is
developmentally normal for his age and compared to other
siblings. He was anthropometrically within normal limits.
The physical examination of the child revealed an
atrophied and shortened left forearm as compared to the
opposite normal limb and he had a single forearm bone
with a curvature laterally. The movements of the left elbow
flexion and extension were normal. All the distal
movements including the rotatory movements of the
forearm and the wrist and the fine finger movements were
possible. The thumb was hypoplastic (Figure 1). The
systemic examination was normal. Evaluation of the
radiographs of both the upper limbs revealed complete
aplasia of the left radius and hypoplasia of the first digit
(thumb) of the left hand, including its metacarpal and the
phalanges. The ulna on the affected side was shorter than
that on the contra lateral side and there was radial deviation
at the wrist (Figure 2). Chest X-ray, echocardiogram,
hemogram including the platelet count, and ultrasound of
the abdomen were normal. Parents were explained the
need of reconstruction surgery and the benefit of
intervention at an early age to which they agreed. He was
operated under general anaesthesia with osteotomy of ulna
at midshaft level with K-wire fixations to centralize the
ulnar bone (Figure 3). Intra operative and post operative
periods were uneventful and he was discharged on day 5
of post operative period.

Figure 1: Short left forearm with hypoplastic thumb
and radial deviation of hand.

Figure 2 (A and B): X-ray of left upper limb of
complete absence of radius and hypoplastic thumb
with radial deviation wrist and ulnar bowing.
DISCUSSION

Our case was unilateral radial club hand type IV with a complete absence of the radius, along with a hypoplastic thumb. Due to the availability of better medical facilities and awareness among parents, most of the radial club hands are diagnosed during the first year of life, with a better functional outcome after reconstructive surgery as in our case. Although radial deficiency can occur in isolation, it is many times associated with other congenital malformations. Forty per cent of the patients with unilateral club hand and 27% with bilateral club hand have associated congenital anomalies involving the cardiac, genitourinary, skeletal, craniofacial and the haematopoietic system. Our case didn’t have any associated congenital anomaly. In Radial club hand, the forearm is foreshortened, with a marked curving of the forearm, stiffness of the elbow and fingers, with the wrist being positioned in radial deviation, and the thumb being either small or absent. The cardiac, genitourinary, skeletal and haematopoietic system involvement requires clinical, radiographic, echocardiogram and laboratory evaluation as appropriate. The commonly associated syndromes include:

Holt Oram syndrome

Radial dysplasia is associated with congenital heart disease (usually ASD or VSD). Abnormalities of the radius can occur in association with heart disease, but do not qualify as the Holt Oram syndrome when they are not bilateral, they lack the carpals and are associated with other visceral malformations and cardiac malformations which are different from intracardiac shunts, conduction disturbances or pulmonary hypertension which are characteristic of the Holt Oram syndrome.

Thrombocytopenia absent radius (TAR) syndrome

It has an autosomal recessive inheritance. The thrombocytopenia is present at birth. It is differentiated from other conditions by the presence of the thumb.

VACTERLS association

Each letter in this syndrome’s name constitutes an acronym for the defects which are involved: vertebral, anal, cardiac, tracheoesophageal, renal, limb and single umbilical artery. Babies who have been diagnosed with the VACTERLS association usually have at least three or more of these individual anomalies.

Cornelia de Lange syndrome

Children who are affected by this syndrome are usually growth retarded and they have microcephaly, classic facial features, micromelia, sensorineural hearing loss, genitourinary abnormalities and behavioural problems.

Fanconi’s anaemia

It is also a rare autosomal recessive disease. In infancy, there are the usual characteristic facial features (microphthalmos, strabismus and hearing defects). Pancytopenia usually does not present until later in childhood, with the mean age of onset being 8 years. There is an increased susceptibility to malignancy, particularly leukaemia.

Other associations include Seckel’s syndrome and an association with trisomies.

Our case had isolated radial club hand and didn’t fit into any of the above-mentioned defined syndromes. Whenever a club hand is identified, it is imperative to conduct a thorough examination and a diagnostic evaluation of the new born to delineate the associated anomalies that may suggest a syndrome. Once the birth defects have been identified, a treatment plan needs to be developed for the infant, with the gastrointestinal, renal and cardiac anomalies usually requiring early surgical management. If the patient survives these surgeries, the prognosis is usually good. The orthopaedic abnormalities can be treated individually. In our case, the child was younger, with deformities in the left upper limb and no contracture at the elbow joint. He was an ideal case to operate on for a better functional outcome. However, documentations are there for successful corrective surgery, even in older children with radial club hand.

CONCLUSION

Whenever a club hand is identified, it is imperative to conduct a thorough examination and evaluation of the new born to delineate the associated anomalies that may suggest a syndrome, because an early diagnosis and appropriate treatment have a better outcome.
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