Bilateral Radial Club Hands at the Komfo Anokye Teaching Hospital, Kumasi, Ghana: A Case Report.

Konadu-Yeboah Dominic¹,², Peter Konadu³, Yifieyeh Abiboye C⁴, Ampem-Darkwa⁵, Zainab Schumacher⁵

¹Orthopaedic and Trauma Surgeon, Komfo Anokye Teaching Hospital, Kumasi, Ghana, Part-Time Lecturer, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana.
²Orthopaedic and trauma Surgeon, Komfo Anokye Teaching Hospital, Kumasi, Lecturer, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana.
³Paediatric surgeon, Komfo Anokye Teaching Hospital, Kumasi Ghana, Part-Time Lecturer, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana.
⁴Medical Officer, Komfo Anokye Teaching Hospital, Kumasi, Ghana
⁵Surgical trainee, Komfo Anokye Teaching Hospital, Kumasi, Ghana

Abstract

Radial Club Hand (RCH) is a rare and complex congenital disorder of the pre-axial border of the upper extremity. It is a congenital longitudinal radial ray deficiency which presents as radial deviation of the hand and distal forearm in the shape of a golf club [1,2]. Radial ray deficiency (the most common type of longitudinal failure of formation) is a spectrum of malformations affecting the structures of the radial side of the forearm, including hypoplasia of the bones, joints, muscles, tendons, ligaments, nerves and blood vessels. The thumb dysfunction, wrist instability and short upper extremity that accompany this condition can cause significant functional impairment particularly in patients with bilateral involvement. Thus, the performance of activities of daily living such as accomplishing personal hygiene, fastening buttons and zippers becomes challenging [3].

Corresponding author: Dominic Konadu-Yeboah, Senior Orthopaedic and Trauma Surgeon, Komfo Anokye Teaching Hospital, Kumasi, Ghana, Part-Time Lecturer, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana, Phone: +233 206 300 663, Email: domiYk@yahoo.com

Keywords: complex congenital disorder, radial ray deficiency, forearm

Accepted: Sep 24, 2019   Published: Sep 28, 2019
Introduction

Radial Club Hand (RCH) is a rare and complex congenital disorder of the pre-axial border of the upper extremity. It is a congenital longitudinal radial ray deficiency which presents as radial deviation of the hand and distal forearm in the shape of a golf club [1,2]. Radial ray deficiency (the most common type of longitudinal failure of formation) is a spectrum of malformation affecting the structures of the radial side of the forearm, including hypoplasia of the bones, joints, muscles, tendons, ligaments, nerves and blood vessels. The thumb dysfunction, wrist instability and short upper extremity that accompany this condition can cause significant functional impairment particularly in patients with bilateral involvement. Thus, the performance of activities of daily living such as accomplishing personal hygiene, fastening buttons and zippers becomes challenging [3].

The incidence of radial ray-deficiency, including patients with hypoplastic thumb alone varies between 1 in 30,000 and 1 in 100,000 live births, with a male to female ratio of 3:2 [4,5].

In humans the upper limbs form between the 4th and the 8th post-ovulatory weeks. The molecular basis of RCH is unknown but the cause is thought to be a defect in the mesodermal development due to a genetic or environmental insult to the apical ectodermal ridge (AER) occurring between the 4th and 5th embryonic week [6,7]. The AER is a thickened layer of ectoderm that directs differentiation of the underlying mesenchymal tissue and limb formation. Defects in the sonic hedgehog signaling pathway have been identified [4,8]. Early prenatal exposure to antiepileptic drugs (valproic acid, phenobarbital), thalidomide and aminopterin has been associated with radial deficiency [9].

Affected individuals have a high incidence of medical and musculoskeletal abnormalities that increases with increasing severity of the deficiency [10]. Table 1 highlights some of these abnormalities. [11, 12, 13, 14, 15].

Radial ray deficiency has a wide range of phenotypic expression from hypoplasia of the thumb to complete absence of the radius and the first ray, as shown in table 2 below. [15]

There have been very few reports of this condition from Africa. We report our experience in the management of a case of bilateral RCH in a boy who was treated surgically at the age of 19 months.

Case Report

A 1 month old infant presented to our hospital with deformed forearms since birth. He was born to a 32-year-old multiparous petty trader at term through a spontaneous vaginal delivery after an uneventful pregnancy. His mother neither ingested alcohol nor smoked cigarette before or during the pregnancy nor suffered from any chronic illness. She used only the

<table>
<thead>
<tr>
<th>Syndrome/Anomaly</th>
<th>Components</th>
</tr>
</thead>
<tbody>
<tr>
<td>VACTERL</td>
<td>Vertebral, Anal, Cardiac, Tracheoesophageal, Renal, Limb</td>
</tr>
<tr>
<td>HOLT-ORAM</td>
<td>Absent radius, Heart defect (Atrial septal defect or heart block)</td>
</tr>
<tr>
<td>FANCONI ANAEMIA</td>
<td>Radial ray deficiency, Pancytopenia</td>
</tr>
<tr>
<td>OKIHIRO</td>
<td>Radial ray deficiency, Duane anomaly (Congenital type of strabismus)</td>
</tr>
<tr>
<td>RAPADILINO</td>
<td>Cleft palate, Patellar aplasia, Radial ray deficiency</td>
</tr>
<tr>
<td>TOWNE-BROOK</td>
<td>ARM, abnormal external ear associated with sensorineural deafness and thumb malformations</td>
</tr>
<tr>
<td>TAR</td>
<td>Thrombocytopenia, absent radius</td>
</tr>
</tbody>
</table>
recommended medicines (folate and ferrous sulphate) during her hospital supervised pregnancy. She, however, had no fetal anomaly scan during this period. No other family member had such a presentation.

Physical examination revealed a young infant who was active. Both forearms were shortened and bowed ulnarwards. There were no abnormalities detected in other systems. X-ray confirmed bilateral Bayne type IV deformity, Figure 1. Complete blood count was normal.

A 6-week serial passive stretching and casting was performed at our physiotherapy department. The initial passive stretching rendered the soft tissues supple to facilitate surgical correction. He was then prepared for operative correction of the anomalies and surgery was performed, figure 2.

Our operative procedure began with a five-centimeter incision along the subcutaneous border of the ulna on both sides. The wrist and forearm deformities of both sides were corrected at the same sitting, we first corrected the deformity on the right side followed by that on the left. Next, a medial based closing wedge osteotomy of the ulna was performed at the mid shaft and the contracted radial wrist soft tissues were released via a five-centimeter transverse incision along the wrist flexion skin crease on the radial aspect of the wrist. This was done with care to preserve the branches of the superficial radial nerve. The flexor carpi radialis tendon was detached routed subcutaneously and attached to the extensor carpi radialis brevis tendon end-to-side using Pulvertaft weaves and nylon 3-0 suture, for extension of the wrist.

The ulnar osteotomy was stabilised with an intramedullary 2.0mm Kirchner wire (K-wire). Wrist bridging external fixators were mounted bilaterally to maintain correction until 12 weeks after surgery. Intraoperative fluoroscopic image of the correction of the left forearm is depicted in figure 3.

The K-wires were removed at six weeks following surgery and active motion of the wrists and digits was encouraged. The external fixators were removed 12 weeks post operatively. Figure 3 demonstrates corrected anomalies after removal of external fixators. He was discharged home with a brace to be applied at night to maintain the new positions of the wrists, after removal of the external fixators, for 12 additional weeks.

There was satisfactory correction of his deformities at his last out-patient review with acceptable mobility of the wrist, metacarpophalangeal and interphalangeal joints on either side with adequate grip strength of both hands. Active and passive physiotherapy are still on-going.

Discussion

The diagnosis of RCH is suspected clinically and confirmed with plain radiographs of the affected limbs. Our patient’s presentation was non-syndromic and there was no identifiable risk factor in his medical history. The management of RCH should commence soon after birth and includes serial manipulation and splinting. The goal is to stretch the contracted soft tissue and radial structures to achieve passive correction of the deformity by aligning the hand and the wrist with the ulna. Sufficient soft-tissue stretching before surgery is a prerequisite for a successful surgery. This should be performed earnestly until the time of surgery [16]. This necessary conservative optimization for our patient lasted 18 months which also allowed for growth of

<table>
<thead>
<tr>
<th>CLASS</th>
<th>DESCRIPTION</th>
<th>ANATOMY</th>
</tr>
</thead>
<tbody>
<tr>
<td>TYPE I</td>
<td>Short distal radius</td>
<td>Radius slightly shorter than ulna (distal growth plate present)</td>
</tr>
<tr>
<td>TYPE II</td>
<td>Hypoplastic radius</td>
<td>Radius is smaller than ulna (no growth plate is present)</td>
</tr>
<tr>
<td>TYPE III</td>
<td>Partial absence of radius</td>
<td>Only a small proximal radius is present</td>
</tr>
<tr>
<td>TYPE IV</td>
<td>Total absence of radius</td>
<td>Radius absent Ulna may be curved</td>
</tr>
</tbody>
</table>

Table 2. Radiological Classification of Radial Ray deficiency by Bayne
Figure 1. Depicts pre-operative radiographs of the forearms of the infant showing bilateral absence of the radius.

Figure 2. Immediate pre- and post-operative pictures with both forearms in supination.
Figure 3. Fluoroscopic image showing the K-wire running from the 3rd metacarpal to bridge the ulnar osteotomy. The Schanze screws of the external fixator are shown, one each in the first metacarpal and in the proximal ulna with the forearm in pronation.

Figure 4. The appearance of the forearms at 12 weeks after removal of the external fixators.
tissues and thus made the subsequent surgical correction easier. Fig 4.

A popular surgical method is the centralisation of the carpus with tendon transfer so as to maintain the corrected position of the carpus [17,18]. The first centralization of the wrist into the distal ulna was performed by Sayre in 1894 [19]. Gradual correction using the Ilizarov method for Bayne Type III or IV has also been described. Severe radial deviation of the wrist recurred [20]. To date there is no established treatment method for mild radial ray deficiency (Bayne Type I or II). In severe cases of radial dysplasia, ulna shortening osteotomy facilitates centralization of the carpus [21]. In this report we present our centralization procedure for Bayne Type IV with ulnar shortening osteotomy, soft tissue release of the radial wrist and maintenance of correction of deformity with a unilateral uniplanar wrist-bridging external fixation.

The post-operative course was uneventful and our outcome satisfactory. The child would be followed for recurrence of radial deformity.

Conclusion

We have presented our management of a boy with Bilateral RCH (Bayne Type Type IV) starting with serial stretching and casting of deformities followed by ulnar osteotomy which enhanced centralization of the carpus. In Bayne Type IV deformity, soft tissue release, distraction and ulnar osteotomy is a good operative method of correction. A long-term follow-up is recommended.

References


Figure 8. Plant derived biomolecules studied in colorectal cancer
Figure 8. Plant derived biomolecules studied in colorectal cancer

(a) 3,4-Di-O-caffeoylquinic acid

(b) 1′-Acetoxychavicol acetate (ACA)

(c) Piperlongumine

(d) Berberine

(e) Anacardic Acid

(f) Sanguinarine

(g) Cardamonin

(h) Celastrol

(i) Deguelin

(j) Diosgenin

(k) Emodin

(l) Embelin

(m) Scutellarin

(n) Gambogenic acid