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The Association of Radial Deficiency with Thumb Hypoplasia

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Background: Congenital longitudinal deficiencies of the radius and thumb are known to be associated with one another; however, the details of their relationship are unknown. The purpose of this study was to determine whether increased severity of radial deficiencies is associated with increased severity of thumb deficiencies and to review the relationship between radial deficiency and reconstructibility of a hypoplastic thumb.

Methods: Radiographs and charts of 227 affected upper extremities of 139 patients with radial longitudinal deficiency were reviewed. The associated thumb deficiency was classified according to a modification of the Blauth and Schneider-Sickert scheme and the radial deficiency was classified according to a modification of the Bayne and Klug criteria for 191 extremities of 119 patients.

Results: The severity of the thumb deficiency was directly proportional to the severity of the radial deficiency (p < 0.0001). Half of the extremities had either a thumb deficiency or thumb and carpal deficiencies without radial deficiency. Two-thirds (sixty-three) of the ninety-five limbs with a normal radius had a thumb that could be surgically reconstructed. Seventy-one (91%) of seventy-eight extremities with a thumb amenable to surgical reconstruction had a radius that did not require surgical reconstruction. All extremities with a radial and/or carpal deficiency had a thumb deficiency. Forty-eight (94%) of fifty-one extremities with complete absence of the radius had a thumb that was not reconstructible.

Conclusions: This study supports the growing body of evidence that the components of radial longitudinal deficiency represent a progressive spectrum of upper extremity abnormalities, and a distal progression of severity, with distal structures likely to be more involved than proximal structures.

Level of Evidence: Prognostic study, Level II-1 (retrospective study). See Instructions to Authors for a complete description of levels of evidence.

Longitudinal deficiency of the radius and thumb hypoplasia are complex congenital disorders that may require surgical treatment to maximize hand function. Although the two abnormalities are known to be associated, few studies of large populations of children with these unusual conditions have been published. The prevalence of deficiency of the radial aspect of the forearm and hand is between 1:30,000 and 1:100,000 live births, and the male:female ratio is 3:2. The prevalence of bilateral involvement is between 64% (sixty-four of 101) and 72% (forty-nine of sixty-eight), and radial deficiency is commonly linked with several syndromes including Holt-Oram syndrome, thrombocytopenia absent radius (TAR) syndrome, Fanconi syndrome, and the VACTERL (vertebral anomalies, anal atresia, cardiac anomalies, tracheoesophageal fistula, renal anomalies, radial anomalies, lung anomalies) association.

We previously studied a large population of patients with thumb deficiency to describe associated anomalies and to determine the prevalence of various types of hypoplastic thumbs according to our modification of the Blauth and Schneider-Sickert classification (Table I and Figs. 1-A through 1-G). This scheme allows the surgeon to classify thumb hypoplasia on the basis of clinical and radiographic findings in order to determine whether the thumb can be reconstructed or whether thumb ablation and index finger pollicization would be the preferred treatment.

In a subsequent study of this same population, we determined that radial deficiency could be classified with a modification of the Bayne and Klug system, which recognizes the spectrum of pathological entities involved in this condition, including radial deficiency, carpal anomalies, and hypoplasia of the thumb. Carpal anomalies are an integral component of radial deficiency and link thumb hypoplasia with radial deficiency. Isolated thumb hypoplasia represents radial deficiency...
in its mildest form, thumb hypoplasia and carpal anomalies represent an intermediate form, and absence or abnormality of all of the radial structures (thumb, radial carpus, and radius) is at the extreme end of the spectrum. This modified classification system is presented in Table II and illustrated in Figures 2-A through 2-D. Use of this classification also assists in treatment decisions; the type-1 radius usually requires no surgical intervention, whereas the type-2 radius may be amenable to lengthening, and the severely deficient radius (type 3 or 4) may benefit from surgical treatment, including centralization (surgical reduction of the carpus onto the distal end of the ulna).

Both of our previous studies suggested an association between the severity of radial deficiency and the severity of thumb hypoplasia. The purpose of the current study was to determine if increased severity of radial deficiency is associated with increased severity of thumb deficiency. We hypothesized that the degree of radial deficiency would correlate directly with the degree of thumb hypoplasia. Because the degree of thumb hypoplasia determines the reconstructibility of the thumb, we also hypothesized that this relationship could be used to predict the reconstructibility of a hypoplastic thumb.

Materials and Methods

The charts and radiographs of all subjects with a diagnosis of deficiency of the radius or thumb seen at Shriners Hospitals for Children, Northern California, between 1923 and 2002 (124 patients) or seen at the private office of one of the authors (H.R.McC. Jr.) between 1978 and 2000 (fifteen...
patients) were reviewed. Our institutional review board approved the retrospective chart and radiograph review. The findings on physical examination, radiographs, and operative findings for these 139 patients with 227 affected upper extremities were studied to determine if adequate clinical, radiographic, and operative information was available to classify the radius and thumb malformations according to the modified Bayne and Klug and Blauth and Schneider-Sickert systems and to determine if there were associated malformations that called into question the diagnosis of radial longitudinal deficiency. Subjects were excluded from the

<table>
<thead>
<tr>
<th>Table I</th>
<th>Modification of Blauth and Schneider-Sickert Classification (Key Features Shaded)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type 1</td>
<td>Type 2</td>
</tr>
<tr>
<td>Thumb size</td>
<td>Normal or small</td>
</tr>
<tr>
<td>First web</td>
<td>Normal size and location</td>
</tr>
<tr>
<td>Intrinsic muscles</td>
<td>APB and OP hypoplasic</td>
</tr>
<tr>
<td>Extrinsic muscles</td>
<td>Normal</td>
</tr>
<tr>
<td>Ligaments</td>
<td>Normal</td>
</tr>
<tr>
<td>Bones and joints</td>
<td>All bones present, maybe hypoplasic</td>
</tr>
</tbody>
</table>

*APB = abductor pollicis brevis, OP = opponens pollicis, FPB = flexor pollicis brevis, FPL = flexor pollicis longus, EPL = extensor pollicis longus, MP UCL = metacarpophalangeal ulnar collateral ligament, and RCL = radial collateral ligament.

Fig. 1-C
Type 3A, which has all of the characteristics of type 2 with the addition of abnormal extrinsic muscles, resulting in limited motion of the interphalangeal joint, as shown here, or other abnormalities such as pollex abductus.

Fig. 1-D
Radiograph of a type-3A thumb deficiency (in a different patient from the one seen in Fig. 1-C). The base of the first metacarpal is small but present, another key feature of type-3A thumbs. This hand also has pollex abductus.
study if inadequate information was available for classification of either the thumb or the radius or if they had an ipsilateral triphalangeal thumb, delta phalanx in the thumb, duplicated thumb, ulnar hypoplasia, aphalangia of fingers, or symbrachydactyly. Six patients (twelve affected extremities) with TAR syndrome were also excluded because the thumb deficiency could not be classified with use of the modified Blauth and Schneider-Sickert criteria, although all of them had an absent radius and a functioning but abnormal thumb. One hundred and nineteen patients with 191 affected limbs fit the inclusion criteria for this study. One hundred and four of these patients had been included in the previous study of the classification of radial deficiency \(^\text{12}\), and ninety-eight had been included in the previous study of thumb deficiency \(^\text{10}\).
Forty-eight (67%) of the seventy-two children with bilateral radial deficiency had asymmetric severity. Twelve associated syndromes were diagnosed in fifty-five patients (46%) (Table III). Bilateral involvement was more common in children with a syndrome; 78% had bilateral deficiency, and 22% had a unilateral anomaly (Table IV). Children with bilateral deficiency were more likely to have a syndrome: 60% had a syndrome compared with 26% of the children with unilateral deficiency (Table IV). The true prevalence of syndromes in this population is probably higher than we detected, as genetic consultation was not available at our institution before 1978. Twenty patients in this series were born before 1960 and thus were ineligible for care at Shriners Hospital by 1978.

Thumb deficiency was classified according to a modification of the scheme described by Blauth and Schneider-Sickert. Classification with this system requires detailed clinical examination and adequate radiographs (Table I and Figs. 1-A through 1-G). Important structures for classification include the size of the first web space, the presence and function of the intrinsic and extrinsic muscles of the thumb, the presence and function of the collateral ligaments of the metacarpophalangeal joint, and the presence and size of the bones of the first ray. Types 1, 2, and 3A thumbs can be reconstructed surgically (although type 1 does not usually require reconstruction). Types 3B, 4, and 5 are treated with ablation of the thumb, if present, and pollicization of the index finger. For six patients, sufficient information was available to determine

<table>
<thead>
<tr>
<th>Type</th>
<th>Thumb</th>
<th>Carpus</th>
<th>Distal Part of Radius</th>
<th>Proximal Part of Radius</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Hypoplastic or absent</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>0</td>
<td>Hypoplastic or absent</td>
<td>Absence, hypoplasia, or coalition</td>
<td>Normal</td>
<td>Normal, radioulnar synostosis, or congenital dislocation of radial head</td>
</tr>
<tr>
<td>1</td>
<td>Hypoplastic or absent</td>
<td>Absence, hypoplasia, or coalition</td>
<td>&gt;2 mm shorter than ulna</td>
<td>Normal, radioulnar synostosis, or congenital dislocation of radial head</td>
</tr>
<tr>
<td>2</td>
<td>Hypoplastic or absent</td>
<td>Absence, hypoplasia, or coalition</td>
<td>Hypoplasia</td>
<td>Hypoplasia</td>
</tr>
<tr>
<td>3</td>
<td>Hypoplastic or absent</td>
<td>Absence, hypoplasia, or coalition</td>
<td>Physis absent</td>
<td>Variable hypoplasia</td>
</tr>
<tr>
<td>4</td>
<td>Hypoplastic or absent</td>
<td>Absence, hypoplasia, or coalition</td>
<td>Absent</td>
<td>Absent</td>
</tr>
</tbody>
</table>

TABLE III Associated Syndromes

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>No. of Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>VACTERL</td>
<td>22</td>
</tr>
<tr>
<td>Holt-Oram</td>
<td>21</td>
</tr>
<tr>
<td>Nager</td>
<td>2</td>
</tr>
<tr>
<td>Juberg-Hayward</td>
<td>2 (identical twins)</td>
</tr>
<tr>
<td>Fanconi anemia</td>
<td>1</td>
</tr>
<tr>
<td>Other</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>55</td>
</tr>
</tbody>
</table>
that they had either a type-2 or type-3A thumb, but there was not enough information to determine between these two types. Thus, for the purposes of statistical analysis, types 2 and 3A were combined (Table V).

Radial longitudinal deficiency was classified according to our modification of the Bayne and Klug criteria. This system is based on radiographic evaluation of both the radius and the carpus (Table II and Figs. 2-A through 2-D). Several carpal anomalies are seen in association with radial and thumb deficiency, including hypoplasia and absence of the scaphoid, trapezium, and lunate as well as carpal coalitions. Normal radiographic ossification of the scaphoid begins by the age of eight years, and therefore only children eight years of age and older at the time of the most recent radiographic examination were classified as having an absent scaphoid (type 0). Children who were seven years of age or younger were classified as having a type-N or 0 deficiency. Since the classification of type-N radii in children who are less than eight years old may change as the children grow older, types N and 0 were combined into one category for the statistical analysis (Table V).

Statistical Analysis
The correlation between thumb deficiency and radial deficiency was analyzed with use of Spearman rho and Kendall tau with significance determined by p < 0.01.

Results
Radial deficiency was highly correlated with thumb deficiency; all extremities with radial or carpal deficiency had thumb deficiency (Table V). Ninety-five (50%) of the 191 extremities had thumb deficiency with no radial deficiency (type N) or thumb and carpal deficiency with no radial deficiency (type 0). Forty-eight (94%) of the fifty-one limbs with complete absence of the radius (type-4 radial deficiency) had a non-reconstructible thumb (type-4 or 5 thumb deficiency), whereas sixty-three (66%) of the ninety-five limbs with a normal radius had a reconstructible thumb (type-1, 2, or 3A thumb deficiency).

In addition, seventy-one (91%) of the seventy-eight extremities with a reconstructible thumb (type 1, 2, or 3A) had a radius that was either of normal length or did not require surgical reconstruction (type N, 0, or 1). Of the 113 extremities with a non-reconstructible thumb (type 3B, 4, or 5), fifty-eight (51%) had a type-2, 3, or 4 radius.

Spearman rho and Kendall tau calculations both showed a highly significant and positive association between the severity of radial deficiency and the severity of thumb deficiency as classified with the modified Blauth and Schneider-Sickert and Bayne and Klug systems, with a p value of <0.0001 for both. The Spearman correlation was 0.6007, and the Kendall correlation was 0.5260.

Discussion
The underlying cause of thumb and radial deficiencies and their true relationship with each other remain unknown. In this clinical series of patients with thumb and radial deficiencies, proximal deficiency (i.e., of the radius) was always associated with distal deficiency, but distal deficiency often occurred in isolation. Several authors have reported the association of thumb deficiency with radial deficiency. Previous descriptions have focused on the association between thumb deficiency and carpal anomalies as well as the association between radial deficiency and carpal anomalies. Published case reports

<table>
<thead>
<tr>
<th>TABLE IV Bilateral and Unilateral Deficiency According to Whether the Patient Had a Syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deficiency (no. of patients)</td>
</tr>
<tr>
<td>Syndrome</td>
</tr>
<tr>
<td>No known syndrome</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TABLE V Occurrence of Radial and Thumb Types*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thumb Type†</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2 or 3A</td>
</tr>
<tr>
<td>3B</td>
</tr>
<tr>
<td>4</td>
</tr>
<tr>
<td>5</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

*Blue = thumb deficiency with no radial deficiency, red = non-reconstructible thumb and absent radius, light blue = reconstructible thumb and normal radius, green = reconstructible thumb and radius that is deficient but does not require reconstruction, and yellow = non-reconstructible thumb and very deficient radius. †Thumb types 2, 3A, and “2 or 3A” have been combined. ‡Radial types N, 0, and “N or 0” have been combined. §Five of these seven extremities belonged to three patients, two with VACTERL association and one with Goldenhar syndrome.
of carpal deficiency have usually described either thumb and/or radial deficiency coexisting with carpal abnormality; isolated carpal absence is extremely rare and did not occur in our series. Patients with TAR syndrome were excluded from our series because the thumb deficiency could not be classified with use of the modified Blauth and Schneider-Sickert classification. These patients all have type-4 radial deficiency and do not appear to have distal progression of severity.

Although the etiology of radial deficiency is unknown, it is certainly multifactorial. Specific genes are known to be associated with limb development, and some cases of radial deficiency are clearly genetic. Associations with multiple congenital anomaly syndromes suggest that radial deficiency may occur with the deletion of chromosome 22q11. Holt-Oram syndrome appears to be heterogeneous, with linkage to gene map locus 12q24.1 in some cohorts and TBX5 gene mutations in others. Fanconi anemia has been mapped to gene locus 16q24.3, and diagnosis with chromosomal breakage analysis has made it possible to treat this previously fatal aplastic anemia with bone marrow transplantation. TAR syndrome has not yet been mapped.

Thus, thumb deficiency is associated with many syndromes mapped to numerous loci throughout the genome. Other heritable thumb malformations, such as triphalangeal thumb and thumb polydactyly, have been mapped to chromosome 7q36.

Fetal exposure to toxins such as thalidomide can cause radial deficiency. It is possible that some cases of radial deficiency are caused by unknown environmental toxins that interrupt longitudinal development of the upper limb when the fetus is exposed to them at a critical phase of development. Unique features of individual digits are known to develop in utero in response to the zone of polarizing activity, a specialized area of the posterior apical epidermal ridge. In the vertebrate limb bud, signals from the zone of polarizing activity are known to control the pattern of cellular differentiation in a radioulnar direction. The zone of polarizing activity, through mechanisms currently under investigation, has some regulatory power over the apical epidermal ridge. Digit formation is also under the control of the zone of polarizing activity, which seems to be responsible for ordering digit type and number and is therefore potentially responsible for forming the unique features of the thumb. As early as 1977, Lamb suggested a localized absence of the apical epidermal ridge as a cause for radial deficiencies. More recently, sonic hedgehog protein (Shh) has been shown to mediate the activity of the
zone of polarizing activity\textsuperscript{46}, and, in mice, disruption of the Shh pathway has been associated with a spectrum of developmental anomalies very similar to VACTERL\textsuperscript{46}.

The interaction of the apical epidermal ridge with the underlying undifferentiated mesoderm is responsible for limb outgrowth in a proximal-to-distal direction, with proximal...
of a spectrum of abnormal upper-extremity development rather than independent processes. The severity of these malformations progresses distally and thus the presence of a severe radial deficiency signals the presence of a severely deficient or absent thumb.

**References**


38. Josefson D. Couple select healthy embryo to provide stem cells for sister. BMJ. 2000;321:917.


