Symbrachydactyly

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Indian J Plast Surg

Abstract

Symbrachydactyly is a rare congenital hand malformation in which a child is born with abnormally short digits that may be webbed, misshaped, or missing, and it is usually a unilateral condition. There is no standardized treatment algorithm for the management of symbrachydactyly. The function of the hand is often not adequate and requires early surgical intervention to restore useful prehension and appearance. This CME article presents a brief review of the embryology, history, classification and clinical presentation, and author’s experience of treating 19 children with symbrachydactyly over 10 years. Creation of thumb web, lengthening of thumb, and creating an opposition post results in prehension of hand with an improved quality of life.

Introduction

Symbrachydactyly is a rare congenital hand difference with an incidence of approximately 1 in 32,000 live births. It was once considered a form of a cleft hand and termed atypical cleft hand due to central ray deficiency, which is usually bilateral and often associated with cleft feet.1

These hands do not represent a terminal failure of formation. It is usually a unilateral pathology, characterized by failure of complete formation of the fingers, and present as rudimentary “nubbins” containing elements of the nail plate, bone and cartilage. Typically, the central digits are absent, the border digits are relatively spared, and syndactyly may be present. The child often presents with short, webbed digits.

There is little evidence to show that symbrachydactyly is due to a genetic defect or passed from one generation to another. It is caused by vascular dysgenesis during fetal development, with interruption of the subclavian artery distal to the internal thoracic artery and a decrease in blood flow in the affected limb before the gestational age of 42 days.

Symbrachydactyly usually occurs as an isolated hand difference. Clinical presentations are varied but always include short fingers. The underlying defect is believed to be mesodermal, which explains the presence of remnants of the distal part of the fingers, namely, ectodermal tissue, pulp, nail folds, and nails. Symbrachydactyly can be associated with Poland syndrome, which is characterized by hypoplasia, or absence, of the sternal head of pectoralis major (►Fig. 1).

Children with symbrachydactyly may have one, or more, of the following:

1. Short fingers in which some bones are smaller than normal or missing. Commonly, the middle phalanges are either hypoplastic or absent.
2. The fingers that are joined together or webbed—syndactyly.
3. Nubbins of skin and soft tissues where the fingers would be.
4. A short hand with gaps, that is, absence of central metacarpals.
5. A short forearm.

DOI https://doi.org/10.1055/s-0041-1734579
ISSN 0970-0358

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Most children with symbrachydactyly have a normal thumb, but the first web contracture is a constant. The hand may not function well, and the bones, muscles, ligaments, and nerves are also often affected.

Radiological Examination

1. Antenatal ultrasonogram—Congenital hand differences are diagnosed more effectively with high-frequency ultrasonogram antenatally.

2. Radiograph—Postnatal X-ray identifies the presence or absence of bones.

3. MRI—Postnatal MRI of finger nubbins will identify the soft-tissue abnormalities.

Classification

The International Federation of Societies for Surgery of the Hand (IFSSH) has recently adopted the Oberg, Manske and Tonkin (OMT) classification system, which is based on embryology and molecular biology. Previously characterized as an undergrowth, in the OMT system, symbrachydactyly is characterized as the failure of formation of the proximal-distal axis, which can involve the entire upper limb or the hand plate only. Blauth classification system for symbrachydactyly includes four phenotypes.2

1. The short-finger type, characterized by the presence of a thumb and four coalesced stiff digits, which may have one or more missing phalanges within these digits.

2. The oligodactylic or “atypical cleft hand” type, characterized by a central aplasia of digits, with thumb and fifth digit present.

3. The monodactylic type, characterized by the presence of a thumb and four associated digital remnants.

4. The peromelic type, characterized by a complete absence of all digits at the metacarpal level.

Yamauchi and Tanabu (Table 1) described a classification that included seven types of symbrachydactyly, based on examination and imaging. This classification does not provide guidance for treatment.3 Foucher modified the Blauth classification to make it more useful in respect of treatment (Table 2).4

Authors Series

Nineteen children with symbrachydactyly underwent surgical management in our institute over a period of 10 years. Eighteen were unilateral and one was bilateral. There were sixteen male and three female children. Of the unilateral cases, seven were on the right side and eleven on the left, indicating left-side dominance. These children usually underwent surgery between the ages of 1 and 3 years. The children were followed-up for a period of 8 to 10 years.

Treatment Goals

The goals of reconstruction are to improve the appearance and function of the hand. Selection of specific treatments for symbrachydactyly depends on the clinical presentation and radiological findings. The outcomes depend on the existing hand differences. The family must be counselled for accepting the outcome.

Nonoperative Management

1. Physiotherapy—performed till the child and the parents can understand the problem and give their consent for the surgery.

a. Stretching of the thumb web.

b. Mobilization of stiff fingers both active and passive.

3. Occupational therapy.

Provision of splints and prosthesis for opposition post. However, prosthesis cannot provide sensation; hence, it is of limited value in children.

Opposition paddles may be helpful in patients with monodactyly, because they provide a surface against which they can pinch (Fig. 2). Targeted muscle reinnervation (TMR) and regenerative peripheral nerve interface (RPNI) may become

| Table 1 Symbrachydactyly types based on radiologic pattern on skeletal reduction |
|---------------------------------|----------------------------------|
| The triphalangia type, characterized by a hand with no missing bones, even though many phalanges (usually the middle phalanx) may be short |
| The biphalangia type, characterized by one or more missing phalanges in one or more digits |
| The monophalangia type, characterized by the presence of a thumb and one or more digits containing only one phalanx |
| The aphalangia type, characterized by the presence of a thumb and digital nubbins with no phalanges |
| The ametacarpia type, characterized by absence of the thumb and digits and absence of one or more metacarpals |
| The acarpia type, characterized by the absence of all digits and thumb and one or more carpal bones |
| The forearm amputation type, characterized by absence of the distal portion of the forearm with small rudimentary nubbins on the amputation stump |
useful to children with the pleomorphic type of symbrachydactyly or transverse arrest at later stage of development.

3D printing of the hand may be useful in preparing light-weight hand prosthesis, which can be customized as the child grows.

**Table 2** Foucher’s classification

<table>
<thead>
<tr>
<th>Type</th>
<th>Features</th>
<th>Thumb</th>
<th>Ulnar digit</th>
<th>Interventions</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>All bones and digits present, brachydactyly and syndactyly</td>
<td>Normal</td>
<td>Bones present, brachydactyly or syndactyly</td>
<td>Syndactyly release</td>
</tr>
<tr>
<td>II A</td>
<td>&gt; 2 fingers. Normal thumb, hypoplastic fingers</td>
<td>Normal</td>
<td>Hypoplastic, syndactyly</td>
<td>Nonvascularized toe phalanx transfers, ablation or stabilization</td>
</tr>
<tr>
<td>II B</td>
<td>Functional border digits, variable central nubbins</td>
<td>Normal</td>
<td>Present, variable hypoplasia and stability</td>
<td>Surgery rarely indicated</td>
</tr>
<tr>
<td>II C</td>
<td>“Spoon hand” thumb conjoined with hypoplastic ulnar digits</td>
<td>Present ± stability</td>
<td>Hypoplastic, clinodactyly</td>
<td>Variable</td>
</tr>
<tr>
<td>III A</td>
<td>Monodactyly</td>
<td>Normal</td>
<td>Absent</td>
<td>Vascularized toe-to-hand transfer</td>
</tr>
<tr>
<td>III B</td>
<td>Monodactyly</td>
<td>Hypoplastic and/or unstable</td>
<td>Absent</td>
<td>Variable, vascularized toe-to-hand transfer, thumb stabilization, thumb lengthening</td>
</tr>
<tr>
<td>IV A</td>
<td>Peromelic, wrist mobility</td>
<td>Absent</td>
<td>Absent</td>
<td>Vascularized toe to hand transfer</td>
</tr>
<tr>
<td>IV B</td>
<td>Peromelic, no wrist mobility</td>
<td>Absent</td>
<td>Absent</td>
<td>Surgery not indicated</td>
</tr>
</tbody>
</table>

**Table 3** Components of a symbrachydactyly reconstruction

<table>
<thead>
<tr>
<th>Excision of rudimentary nubbins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formation or preservation of a mobile, unscarred thumb with stable joints</td>
</tr>
<tr>
<td>Formation or preservation of a broad first web space</td>
</tr>
<tr>
<td>Maintenance of a thumb and at least one digit with sensation</td>
</tr>
<tr>
<td>Formation or preservation of an intact fifth ray with MCP joint motion and stability</td>
</tr>
<tr>
<td>Separation of digital syndactylies</td>
</tr>
<tr>
<td>Preservation of digital proximal interphalangeal joint motion in triphalangeal digits and MCP joint motion in biphalangeal digits</td>
</tr>
</tbody>
</table>

Abbreviations: MCP, metacarpophalangeal.

**Surgical Interventions**

Symbrachydactyly is a spectrum of deficiencies from short, or absent, middle phalanges through absent metacarpals to absence of the entire hand. Treatment is variable and depends on the degree of digital suppression.

The aim of surgical treatment is toward improving function and aesthetics. The various surgical components are enlisted in **Table 3**. Reconstruction is oriented to achieve basic requirements for manipulation of objects such as creation of a radial post, an ulnar post, and an adequate web space.

Stability of the posts should be achieved and at least one of them should be mobile. Once this target is achieved, further enhancement is planned by separating the digits, increasing length, and improving mobility.

It is not possible to draw a surgical algorithm for the management of symbrachydactyly, since planning has to be customized, according to availability of local tissues and the extent of deficiency. The presence of transverse bones and abnormal articulations also influence the planning.
The types of presentations, surgical procedures, and the outcome obtained in our clinical experience has been tabulated in Table 4. It is also not possible to compare the functional outcome of various procedures, because of the absent phalanges, varying degree of hypoplasia, and limited mobility of existing digits.

Different surgical interventions planned in the management of symbrachydactyly are listed below. Planning and sequencing of surgeries have to be individualized, according to the need of the child.

1. Thumb web contracture release and lengthening of the thumb.
2. Creation of an opposition or ulnar post.
3. Release of syndactyly between the short fingers.
4. Lengthening of brachydactyly.
5. Free phalangeal transfer with digital augmentation manoplasty.
6. Microvascular toe transfer.

**First Web Contracture Release and Lengthening of the Thumb**

In symbrachydactyly, first web contractures are treated to improve the grasp span, appearance, and function of the thumb. When all the metacarpals are present and the phalanges are not present or replaced with nubbins, the simplest option is to deepen the first web and lengthen the thumb to create some form of crude pinch against the palm or an available phalanx or nubbins.

We have used various surgical procedures to release the first web.

1. Multiple Z-plasties.
2. Dorsal rotation flap.
4. Square flap.
5. Omega flap.

**Z-Plasties**

Multiple Z-plasties in sequence will achieve the maximum release, but lengthening of the thumb is inadequate. It is often used to lengthen the short thumb by deepening the thumb web.

**Dorsal Rotation Flaps**

When there is tightness of the adductor pollicis muscle in addition to skin contracture, a dorsal rotation flap is planned. Through the same skin incision, the adductor pollicis insertion is released and the flap is rotated into the webspace. The donor defect can either be closed primarily or, occasionally, the skin is grafted (Fig. 3A-E).

The dorsal rotation flap can only be planned when all the metacarpals are present, and the dorsum has adequate width. In the absence of few metacarpals, the dorsal width is narrowed, and a rotation flap cannot be performed.

**Groin Flaps**

Symbrachydactyly with a narrow first web and no available local tissue is reconstructed with a distant flap, such as groin flap, following web release. A vertical skin incision is used to separate the first and second metacarpals and release the insertion of the adductor pollicis muscle. The groin flap is an exceptionally effective procedure under these circumstances (Fig. 4A, B, C). When there is a complex musculoskeletal contracture, such as present of transverse bone, abnormal articulations in carpometacarpal joint, or tight adductor muscles, release and groin flap will be the once choice. The disadvantage is the bulkiness of the flap.

**Square Flaps**

The square flap, introduced by Prof. Hiko Hyakusoku, is a better option for release of first web and simultaneous lengthening of the thumb (Fig. 5A, B). The release of web contracture will be nearly twice that of a single Z-plasty, resulting in better widening of the thumb web and lengthening of the thumb. It incorporates an advancing square, which breaks the linear contracture band, with two triangle flaps rotating from the opposite side into the lateral defects (Video 1).

**Video 1**


In some of these children, the thumb lays in the same plane as the rest of the fingers. These children will require rotational osteotomy of the first metacarpal to bring thumb into opposition and pronation. We have performed this...
<table>
<thead>
<tr>
<th>PT. ID</th>
<th>Symbrachydactyly subtype</th>
<th>Sex and age at presentation</th>
<th>Surgical intervention</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Oligodactyly Right hand</td>
<td>F/1 year</td>
<td>Thumb web release by dorsal transposition flap</td>
<td>Adequate web release obtained</td>
</tr>
<tr>
<td></td>
<td>Narrow thumb web</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>Short-finger type Left hand</td>
<td>M/10 years</td>
<td>Distraction Osteogenesis of proximal phalanges of all fingers</td>
<td>Lengthening of fingers was useful to improve function</td>
</tr>
<tr>
<td></td>
<td>Thumb normal All fingers short with absent middle phalanges</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>Oligodactyly Left hand</td>
<td>M/2 years</td>
<td>Syndactyly release —ring and little fingers—omega flap</td>
<td>Tripod pinch achieved</td>
</tr>
<tr>
<td></td>
<td>Thumb normal Index and mid absent</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>Peromelic Left hand</td>
<td>M/1 year</td>
<td>Free phalangeal transfer to stabilize unstable thumb</td>
<td>Child uses the stable thumb to oppose palm to hold objects. Will require double toe transfer to enhance prehension</td>
</tr>
<tr>
<td></td>
<td>Unstable thumb Absent fingers</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>Oligodactyly type Narrow thumb web Right hand</td>
<td>F/1 year</td>
<td>Thumb web reconstruction with a dorsal transposition flap</td>
<td>Adequate thumb web release obtained</td>
</tr>
<tr>
<td>F</td>
<td>Symbrachydactyly Absent thumb web with transversely placed second metacarpal Absent index Left hand</td>
<td>M/2 years</td>
<td>Thumb web release Excision of transverse metacarpal Groin flap cover</td>
<td>Opposition achieved between released thumb and syndactylized ulnar post</td>
</tr>
<tr>
<td>G</td>
<td>Peromelic type Left hand</td>
<td>M/1 year 1. Free phalangeal transfer for little finger nubbin—15 months 2. Toe-to-hand transfer for thumb reconstruction—4 years</td>
<td>Child can perform hook grip using the reconstructed digits and manipulate objects</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Absent thumb Finger nubbins</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>Oligodactyly Narrow thumb web Left hand</td>
<td>M/2 years</td>
<td>Thumb web release by square flap technique</td>
<td>Adequate release of thumb web achieved</td>
</tr>
<tr>
<td>I</td>
<td>Oligodactyly Narrow thumb web Left hand</td>
<td>M/1 year</td>
<td>Thumb web reconstruction with omega flap technique</td>
<td>Adequate thumb web obtained. Child will need further procedures to release finger syndactyly</td>
</tr>
<tr>
<td>J</td>
<td>Oligodactyly type Inadequate thumb web Absent index Right hand</td>
<td>M/9 months 1. Syndactyly release little and ring using omega flap technique + square flap for thumb web deepening 2. Syndactyly release middle and ring fingers using omega flap</td>
<td>Adequate thumb web and three useful digits achieved after reconstruction. Able to use hand with tripod pinch</td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>Oligodactyly Bilateral – Thumb and thumb web normal Rest of the fingers syndactylized</td>
<td>F/3 months 1. Syndactyly release of marginal fingers—omega flap technique—9 months—both hands 2. Syndactyly release of index and mid—both hands—omega flap technique—2 years</td>
<td>Adequate release of all fingers achieved</td>
<td></td>
</tr>
<tr>
<td>L</td>
<td>Oligodactyly Right hand</td>
<td>M/1 year 1. Syndactyly release—ring and little—1 year—omega flap Thumb web deepening by Z-plasty 2. Syndactyly release middle and ring finger—2 years Omega flap</td>
<td>Tripod pinch obtained</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Short thumb Index absent</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>Oligodactyly Narrow thumb web Left hand</td>
<td>M/1.5 years</td>
<td>Thumb web release by square flap technique</td>
<td>Thumb web release adequate</td>
</tr>
</tbody>
</table>

(continued)
procedure in one child with monodactyly form of symbrachydactyly with short thumb.

**Omega Flaps**

The omega flap is another useful tool for release of the first web and, also, other webbed fingers. It has two components.

1. A truncated dorsal flap with a triangular tip at the level of the proximal interphalangeal (PIP) joint.
2. An anchor incision on the volar side, raising two triangular palmar flaps to cover the lateral surfaces of the base of the finger. This technique avoids web creep (Fig. 6A, B, C). Omega flap is a useful tool when the thumb is syndactylized to index, but only at skin level.

**Creation of an Opposition or Ulnar Post**

The creation of an opposition post is particularly important. Usually, the little finger is short with the absence of the middle phalanx. To create an ulnar post, a free phalangeal bone transfer can be used to lengthen the existing little finger. When there is aplasia of the digit, the only possible surgical option is a free vascularized toe transfer. When the thumb is short, longer toe in place of the little finger will help with opposition. The ideal timing of surgery is at 2 to 3 years of age to allow for a reasonable size of vessel for microvascular anastomosis.

**Syndactyly Release**

Syndactyly release between the short digits is essential. It is done in stages. Index and little fingers need to be released early to avoid shortening of digits. Marginal fingers can be released as early as 6 months and the subsequent fingers released 6 months later.

When we have a child with first web contracture, syndactyly of the ulnar three fingers and an absent index finger at the metacarpophalangeal (MCP) joint level, staged reconstruction is planned. At the first stage, the first web is released along with the release of the syndactyly between the ring and little fingers. At the second procedure, 6 months

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**Table 4 (continued)**

<table>
<thead>
<tr>
<th>PT. ID</th>
<th>Symbrachydactyly subtype</th>
<th>Sex and age at presentation</th>
<th>Surgical intervention</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Short-finger type</td>
<td>M/6 months</td>
<td>1. Syndactyly release between index and middle fingers—9 months 2. Syndactyly release between middle and ring fingers—12 months 3. Distraction osteogenesis of proximal phalanges of index, middle and ring fingers to lengthen the fingers</td>
<td>Achieved stable long fingers of equal length. Improvement in appearance and function achieved. Child is now able to use computer keyboard fluently</td>
</tr>
<tr>
<td>O</td>
<td>Oligodactyly</td>
<td>M/2 years</td>
<td>Thumb web release by square flap</td>
<td>Adequate web release obtained</td>
</tr>
<tr>
<td>P</td>
<td>Hole type of Symbrachydactyly</td>
<td>M/1.5 years</td>
<td>Thumb web release; adductor muscle release Groin flap cover</td>
<td>Child was able to oppose the released thumb and touch the remaining fingers. Able to hold objects and manipulate</td>
</tr>
<tr>
<td>Q</td>
<td>Peromelic type</td>
<td>M/6 months</td>
<td>Free phalangeal transfers to stabilize thumb and to stabilize ring and little finger nubbins</td>
<td>Child was able to oppose the stable thumb to the stable digits and hold objects</td>
</tr>
<tr>
<td>R</td>
<td>Peromelic type</td>
<td>M/9 months</td>
<td>Free phalangeal transfer to middle and ring finger nubbins</td>
<td>Child was able to use Stable fingers for manipulation of objects</td>
</tr>
<tr>
<td>S</td>
<td>Monodactyly</td>
<td>M/1 year</td>
<td>1. Thumb web release and thumb lengthening by square flap method 2. Rotation osteotomy of 1st metacarpal to bring the thumb into opposition and pronation 3. Second toe transfer to reconstruct ulnar post</td>
<td>Child was able to manipulate objects using hook grip. Able to hold a pen and write</td>
</tr>
</tbody>
</table>
later, the syndactyly between the middle and ring fingers is released (►Fig. 7A-D). In most of these cases, the middle phalanx is absent, or there is a very short middle finger with an intact terminal phalanx and nail complex. The postoperative video illustrates that good function has been achieved with the reconstructed hand (Video 2).

**Video 2**


**Lengthening of Brachydactyly**

Distraction osteogenesis is useful to lengthen short digits.\(^7\) It can also be used at a second stage after a toe phalangeal transfer to lengthen the transferred toe phalangeal bone.

When the thumb is short, osteotomy of first metacarpal can be performed and lengthening can be achieved by distraction. Metacarpal lengthening can provide more reliable and stable bone formation than bone grafting.\(^8\) A web deepening procedure is often necessary after the distractor is removed and callus is consolidated.

When multiple fingers are short, distraction of proximal phalanges is performed to obtain finger length. A single distractor can be applied to multiple fingers, and simultaneous lengthening of all fingers can be achieved. Placement of pins should be done under C-arm guidance, and injury to epiphysis and articular surfaces should be avoided.

We had a child with symbrachydactyly of short-finger type. This child had a well-developed thumb and little finger. The middle three fingers were short and syndactylized. At the ages of 9 and 12 months, syndactylies of the middle three fingers were released in two stages, and the fingers allowed to grow individually. At the age of 6 years, the child had short and thin proximal phalanges and terminal phalanges, with intact nail complexes in the middle three fingers. The middle phalanges were absent. The child underwent distraction osteogenesis of the proximal phalanges of all three fingers, using a combined distraction device. Although the distraction takes longer, it is said to be more effective than bone grafting with respect of the bone lengthening achieved.\(^9\) We were able to create a gap in the proximal phalanx, which was replaced by auto-osteoynthesis. At the age of 11 years, the child had fingers of a uniform length (►Fig. 8A, B, C). Fig. 8 (A) Child with short-finger type of symbrachydactyly: first stage reconstruction done with syndactyly release. (B) Lengthening of the index, middle, and ring fingers done with distraction osteogenesis. (C) The postoperative appearance of the lengthened fingers.

Because of reduced usage of hand, the phalanges are often osteopenic and hypoplastic. Distraction must be carried in a slow, careful, and supervised manner for a prolonged period to obtain a stable callus. Even after removal of distractor, intramedullary K wires may be required for keeping the callus in alignment for few weeks. Few children might require bone grafting for nonunion of the distracted callus.\(^10\)

**Free Nonvascularized Phalangeal Transfer**

Single or double free nonvascularized toe phalangeal transfer is used in children with symbrachydactyly. To obtain normal growth of the phalanges, the procedure should be done before 18 months of age. The growth plates may remain open to the age of 7 years. This procedure is only appropriate when the soft-tissue envelope is sufficient. It is necessary to harvest the whole phalanx with an intact periosteal envelope to avoid resorption.\(^11\)
Free nonvascularized phalangeal transfer for symbrachydactyly will have a poor outcome if there is inadequate space in the digital sac. If the transferred phalanx is shortened to fit into the digital sac, it will be reabsorbed, as the medullary cavity is exposed. If there is disruption of the periosteal covering, the phalangeal graft will not survive. Donor digit shortening, instability, and deformity can occur after toe phalangeal harvest. To avoid donor site instability, the flexor and extensor tendons should be sutured together in the resultant gap. This avoids a floppy digit, prevents deformity after phalangeal harvest, and minimizes shortening.

Fig. 5 (A) Child with oligodactyly along with a narrow thumb web—Illustration of the planning for a square flap to reconstruct the thumb web and lengthen the thumb. (B) Postoperative view result showing a longer thumb and ability to hold a pen using pincer grip between thumb and little finger. (C) Schematic representation of the square flap procedure. A square flap is planned on one side of the web and two triangular flaps are planned on the other side of the web. The first triangle has an angle of 60° and the second triangle has an angle of 90°.

Fig. 6 A child with symbrachydactyly along with a narrow thumb web, illustrating the planning for release and reconstruction with an omega flap. (A) Volar view. (B) Dorsal view showing the truncated dorsal flap. (C) Postoperative view showing an adequate first web.

Fig. 7 (A) A child with symbrachydactyly along with fusion of all digits. The thumb is normal. The thumb web is contracted. (B) At the first procedure, the little finger was separated from the ring finger by the omega flap technique and the first web released using a square flap. (C) The markings for the second stage to release the syndactyly of the middle and ring fingers by the omega technique. (D) The result showing four independent useful fingers.
When the metacarpals are of equal length with a good soft-tissue envelope, free phalangeal transfers remain quite stable and can achieve up to 90 degrees of motion at the new MCP joint. However, with metacarpals of uneven lengths, direct placement of the free phalangeal transfers onto the metacarpal heads will often lead to angulated deformity and subluxation.

Digital instability was not noticed in most of these patients, because the flexor and extensor tendons were fused by the palmar aponeurosis over the metacarpal heads. While reconstructing the digit, the tendons must be divided and sutured to the palmar and dorsal sides of the transferred phalanx to act as flexor and extensor. This will provide stability to the reconstructed finger. To avoid instability of the digit, the transverse metatarsal ligament should never be disturbed.

A combination of toe phalangeal transfer, web deepening, and distraction lengthening of the transferred phalanges enhance the hand function, provide stable digits, and enable wider grip span in oligodactylyous type of symbrachydactyly.14

We had a child with peromelic type of symbrachydactyly who presented at the age of 1 year with an unstable thumb at the MCP level. He had a remnant of the terminal phalanx with a nail complex. A free nonvascularized phalangeal transfer was performed to stabilize the thumb at the MCP joint level. Reviewing the patient after 8 years, we found that the patient had retained a stable thumb, but it was short. However, he could oppose to the palm with the short thumb and ride a bicycle to his satisfaction and that of his parents (Fig. 9A, B and C).

Fig. 9 (A) A child with peromelic type of symbrachydactyly: The thumb was attached by a small pedicle but unstable; X-ray showing poor development of the phalanges. (B) Dorsal view of a free toe phalangeal transfer at 15 months, stabilizing the thumb. (C) X-ray at 3-year follow-up showing good survival of the transferred phalanx with no bony absorption. (D) At 8 years of age, although the boy has got no fingers in the opposing post, he opposes with the thumb to the palm and is able to hold the handlebar of a bicycle to ride to school.

**Toe Transfer**

Vascularized toe transfer is an ideal treatment for reconstruction of absent radial post or ulnar post. Toe transfer helps in restoration of pinch, grasp, and prehension in monodactylyous form of symbrachydactyly. The success rate of microvascular toe-to-hand transfer by experienced microvascular surgeons is generally more than 95%.15 The transferred digit often regains adequate function and sensation and helps the hand perform self-care.16
Planning has to be done meticulously, since proximal abnormalities like absent or hypoplastic tendons, abnormal vascular pattern and branching and skeletal hypoplasia are more common in congenital hand differences. Proximal nerves can also be absent and may need nerve transfer from adjacent nerves to provide sensation to transferred digit.

Our experience with second toe transfer to replace the little finger as an opposition post is limited. In children with the monodactylous form of symbrachydactyly, addition of digits by free vascularized double toe transfer, will achieve tripod grip, as opposed to the hook grip of a single toe transfer to the little finger (Fig. 10).

Restoration of pinch in adactylous form of symbrachydactyly needs to be performed in two stages. In the first stage, radial postreconstruction is done by second toe transfer. In the second stage, ulnar postrecreation is done by double toe transfer. Growth rate, functional status, and sensory recovery are often satisfactory. Joint motion restoration will be limited and may require secondary surgeries. Extension deficit in PIP joint level and limited mobility of distal interphalangeal joint may be benefited by tenolysis or arthrolysis.

Conclusions

Making the correct diagnosis is the essential first step of managing congenital hand differences. This is especially more appropriate in symbrachydactyly, because of its diverse morphologic presentations.

Surgical treatment of children with symbrachydactyly is challenging but can provide excellent functional outcomes if surgery is performed early. More severe forms are taxing to the ingenuity of the surgeon. Management has to be individualized in each and every child, depending upon the clinical presentations, availability of digits, stability and their mobility.

We have formulated a management protocol (Fig. 11) for these children with symbrachydactyly. This is only a rough outline and guidance toward planning. Children often require a combination of these procedures or additional newer techniques designed, as per the vision and visualization of the reconstructive surgeon.

Sometimes, surgery has little to offer. Despite the adverse appearance of the hand, children will adapt and learn to use it in a way, which might have been considered impossible at
birth. In our experience, it is possible to satisfy parents with the functional outcome for their wards. The children, when they become adults, will be mature enough to use the corrected hand usefully.

Conflict of Interest
None declared.

Acknowledgment
We would like to acknowledge the contribution of Dr. Kavya Somesh in data acquisition, scientific analysis, preparation of the manuscript, and submission of the article. Without her sincere efforts, hard work, intelligent interventions, enthusiastic support, and exemplary motivation, preparation of this manuscript would have been impossible.

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20 Koskimies E, Lindfors N, Gissler M, Peltonen J, Nietosvaara Y. Congenital upper limb deficiencies and associated malformations in Finland: a population-based study. J Hand Surg Am 2011;36(6):1058–1065Fig. 11 Management protocol for symbrachydactyly. Fig. 11 Management protocol for symbrachydactyly.