INTRODUCTION

Terminal transverse amputations are characterized by the absence of distal portions of the extremities extending across the width of the limb/autopod. Hand amputation most often has traumatic etiology and its congenital presentation is very rare. Most of the cases have traumatic etiology and congenital presentation is less common. A series of six individuals with transverse deficiency through the hands is presented in this communication. The cases were congenital, morphologically similar and showed loss of four fingers, most often postaxial. The affected arm was reduced in size compared to the contralateral limb and there was distortion of palmer creases. All cases were sporadic and non-syndromic in nature. The characteristics of these cases were concordant with the symbrachydactyly type III or monodactylous type, when classified according to the scheme proposed by Blauth and Gekeler (1973). The malformation resulted in permanent quality-of-life impairment in these subjects and warrant prosthetic management. Detailed physical and phenotypic features of the patients have been presented.

KEY WORDS: Limb amputations, transverse defects, terminal deficiency, symbrachydactyly, monodactyly, Pakistani subject.
amputations of variable degrees which resulted in loss of four fingers, mostly postaxial (Table-I). The individuals also exhibited affected limb-length discrepancy compared to the contralateral arm. Palmer creases were distorted. In five of the six cases the feet were unremarkable. All individuals had normal IQ and no associated anomaly was evident in gross physical examination. The patients had functional restrictions in their daily/occupational lives. The snapshot of CTA phenotype is given below:

**Case I:** The individual was fourth in the sibship of four. Reportedly, the first pregnancy of his mother was delayed for three years and the first two sibs (females) died in postpartum. The individual was observed to have CTA through his left palm. Fingers 2-5 were represented by bead-like remnants and there was distortion of palmer dermatoglyphics (Fig.1A). Roentgenographs revealed aplastic/hypoplastic carpals, absent metacarpals, terminal hypoplasia of first digital ray, and mild shortening of radius/ulna (Fig.1 B-C).

**Case II:** This male patient was observed to have CTA of right hand through the palm. A short thumb was evident in the affected hand (Fig.1D). Roentgenographs depicted absence of several carpals and metacarpals; metacarpal 5 was represented by a small peg-like osseous element, and there was terminal symphalangism of first digital ray (Fig.1E). In the left hand, there was medial inclination of index finger and crowding of carpals. In the feet, there was bilateral hypertrophy of first digital ray with hallux valgus (Fig.1F).

**Case III:** There was CTA of right hand which culminated in four nubbin-like digits and a relatively normal thumb (Fig.2A). Characteristic dermatoglyphics were evident.

**Case IV:** The individual had CTA through the medial axis of left palm and digits 2-5 were completely omitted. The thumb appeared unaffected (Fig.2B).

| Variable         | Individual | I     | II    | III   | IV    | V     | VI    |
|------------------|------------|-------|-------|-------|-------|-------|-------|
| **Demographics** |            |       |       |       |       |       |       |
| Gender/age(yrs)  | M/7        | M/20  | F/8   | F/16  | M/12  | M/8   |
| Geographic origin| Southern-Punjab | Interior-Sindh | Upper-Punjab | South-KPK | North-KPK | North-KPK |
| Caste/language    | Arain/ Punjabi | Lashari/ Saraiki | Pathan/Pushto | Khowar/ Swati/Pushto | |
| Parental consanguinity | Distantly related | First cousin | Distantly related | Non-related | Non-related | Non-related |
| Paternal and maternal age at patient’s birth (year) | 40/38 | 20/18 | 29/22 | 27/23 | 29/28 | 37/30 |
| Patient’s parity | 4 of 4 | 1 of 5 | 1 of 3 | 1 of 7 | 3 of 4 | 7 of 9 |
| No. of normal sibs (B:S) | 0:1 | 1:3 | 1:1 | 4:2 | 1:2 | 3:5 |
| **Phenotype**    |            |       |       |       |       |       |       |
| Affected hand    | Left       | Right | Right | Left  | Left  | Left  |
| Amputation axis  | Palm, median | Palm, proximal | Palm, median | Palm, median | Palm, proximal | Palm, proximal |
| Fingers          | Bead like remnants of fingers 2-5 | Digits 2-5 absent | Bead like remnants of fingers 2-5; Terminal hypoplasia | Digits 2-5 absent | Terminal hypoplasia | Absent |
| Thumb            | Terminal hypoplasia, short nail | Short, distal symphalangism | Terminal hypoplasia | Unaffected | Absent | Terminal hypoplasia, short nail |
| Affected arm, reduced/short | ++ | + | ++ | + | + | No |
| Contralateral arm| Mild shortening of zeugopod and stylopod | Medial inclination of index finger; crowding of carpals | Left thumb with extra palmer creases | Unaffected | Unaffected | Unaffected |
| Others           | Carpals absent; hypoplastic metacarpals | Fused carpals; metacarpals 2-4 not visible; reduced metacarpal 5 | Swelling on left throat | |

+=mild; +++=moderate.
Case V: The left hand exhibited CTA through the proximal plan of palm (Fig.2C). Fingers 1-4 were completely omitted and only the 5th finger was present. The affected arm was markedly reduced in size.

Case VI: The individual had CTA through the left hand (Fig.2D). Fingers 2-5 were amputated at their bases while the thumb showed terminal deficiency.

Fig.1: (A-C): Photographs and roentgenographs of individual I. (D-F): Phenotype in individual II.

Fig.2: (A): Amputation in individual III; (B): individual IV; (C): individual V. (D): individual VI.
DISCUSSION

The CTA is generally reported as symbrachydactyly. In a retrospective study on patients with terminal amputations, Kallemeier et al. concluded that transverse deficiency through the forearm represents a proximal continuum of symbrachydactyly. The six cases presented here show remarkable similarities with each other and were concordant with symbrachydactyly type III or monodactylous type; the hallmark of this type is the absence of all fingers other than the thumb, including parts of the metacarpals. Interestingly, in one of our cases the amputated fingers were preaxial including the thumb and only the 5th finger was present.

Nubbin-like digits appear to be an occasional feature of symbrachydactyly. Two of our patients also exhibited soft nubbin-like finger remnants at the distal border of affected hand. Kallemeier et al. observed that 71% of the 291 patients with upper-extremity transverse deficiency had soft tissue nubbins at the end of their amputations. We also observed that left hand was more commonly affected than the right. De Smet et al. recruited patients with symbrachydactyly and observed that the involvement of left hand was twice as common as right. The authors also witnessed that unilateral cases were customary.

The unilateral and isolated nature in most of the cases of symbrachydactyly supports the nongenetic etiology. However, involvement of this anomaly with other syndromes likes Adams-Oliver syndrome (OMIM-100300) and ADAM complex (OMIM-217100) may suggest genetic factors in certain types. Other hypothesis than the vascular disruption explaining symbrachydactyly and associated syndromes have been proposed. The mesenchymal failure of the arm bud may cause terminal aplasia or intercalated deficiency.

Hand amputations have devastating effect on the lives of the individuals. CTA have not gained much attention in Pakistan. CTA not only affect the functioning of hand but also put great social and psychological burden on the patients. Such individuals remain highly disadvantaged in daily and occupational lives. There is a dire need to launch further studies for a comprehensive understanding of the prevalence, determinants and etiology of these anomalies in Pakistan.

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Authors’ Contribution:
SM conceived and designed the study.
HFR, KL, SU, NAB and WU did the field work and collected data.
SM drafted the manuscript. All authors read and approved the manuscript.