Unusual Type of Hand Preaxial Polydactyly

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Introduction

Polydactyly is one of the frequently observed congenital anomalies of the hand and foot. In this anomaly, a rudimentary or fully developed additional digit is seen along with the usual digits. The anomaly is also referred as polydactylia, polydactylysm, or hyperdactyly.1 It can occur as a part of a syndrome or isolatedly.2 Polydactyly of the hand or of the foot are categorized into preaxial, postaxial and central types.3 Thumb polydactyly is the most common type of polydactyly in the hand. Currently, the classification of thumb polydactyly proposed by Wassel4 is in clinical use. In the present case, we report an unusual type of polydactyly and discuss its structural components.

Case Report

During regular dissections for medical graduates, we came across a rare type of preaxial polydactyly in the left hand. It was observed in an ~55-year-old formalin embalmed male cadaver of South Indian origin. The additional digit was present on the lateral margin (preaxial border) of the thumb. The extra digit was ~1.4 cm in length, and it was found to be budding at the level of the base of the distal phalanx (∆ Fig. 1 and 2). A dissection was performed to evaluate the structural framework of the digit. Upon the reflection of the skin, a mass of fatty tissue was observed (∆ Fig. 3). Upon further removal of the fatty tissue, a thick cord of connective tissue was noted in its central core. This cord was surrounded by a mass of fatty tissue. The skeletal, muscle and tendon components were found to be absent in the additional digit.

Discussion

The pathogenesis of preaxial polydactyly has been studied in both experimental animals5,6 and in men.7 Yasuda has studied 13 human embryos and demonstrated the morphological features in the early pathogenesis of preaxial polydactyly.7 In this study, it is concluded that the interaction between the ectoderm and the mesoderm of the limb plays a crucial role in the development of preaxial
polydactyly. In both white and black populations, the incidence of thumb polydactyly is found to occur in 8 in 100,000 individuals. The Wassel classification is proposed based on the level of bifurcation or duplication, and is widely accepted clinically. According to Wassel, the pathoanatomy of thumb polydactyly can be divided into 7 groups; type I: bifid distal phalanx; type II: duplicated distal phalanx; type III: bifid proximal phalanx; type IV: duplication of the proximal phalanx that rests on the broad metacarpal; type V: bifid metacarpal; type VI: duplicated metacarpal; and type VII: triphalangism. The occurrence of type IV is more common when compared with other types. In the present case, we report an unusual type of thumb polydactyly in which the additional thumb was found to be devoid of skeletal, muscle and tendon components. In the Wassel classification of thumb polydactyly, the reported type has not been described. The documentation of this rare type may be clinically important. In an immature hand, due to the presence of the cartilaginous epiphysis, the identification of the level of bifurcation between the duplicated components is difficult. This can possibly explain why sometimes the clinically diagnosed type of polydactyly is different from the findings.
encountered in the surgery and, in such cases, the classification may not be appropriate to deal with. A radiographic examination is usually recommended to evaluate the presence of the skeletal component in the additional digit, before planning the surgery. For the better reconstruction outcome and to avoid or reduce complications in operations, the components of the additional digit such as skin, nail, bone, and the ligaments should be evaluated and treated in a simultaneous manner. The additional digit observed in the present case was non-functional and there were no skeletal, muscle and tendon components in it. The presence of a thick cord of connective tissue formed by the fibrous flexor sheath of the flexor pollicis longus tendon reported here may be clinically significant during the diagnosis, treatment and while planning the surgery and reconstruction procedures.

**Conflicts of Interest**
The authors have no conflicts of interest to declare.

**References**