CAMPTODACTYLY AND KIRNER’S DEFORMITY IN ONE FAMILY

BİR AİLEDEKİ KAMPTODAKTİLİ VE KİRNER DEFORMİTESİ

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ÖZET

Anahtar Kelimeler: Kamptodaktyly, Kirner deformitesi, genetik geçiş

SUMMARY
Kirner’s deformity and camptodactyly are rare congenital hand deformities usually occurring independently. Kirner’s deformity is characterized by palmar and radial deviation of the distal phalanx of the little finger with a varying degree of rotation and a small, dysmorphic watch-glass nail associated with swelling of the distal interphalangeal joint. Camptodactyly is a non-traumatic flexion contracture of the proximal interphalangeal joints of one or more fingers, resulting in a permanent volar inclination of the affected digits. Little finger is the most frequently affected one, but additional fingers may also be involved. The combination of these two entities in a family had not been previously reported. The family presented here suggests that Kirner’s deformity and camptodactyly may have a common genetic factor.

Key words: Kamptodactyly, Kirner’s deformity, genetic transmission

Introduction
Kirner’s deformity and camptodactyly are rare and usually bilaterally occurring congenital hand deformities ¹-³. Kirner’s deformity is characterized by palmar and radial deviation of the distal phalanx of the little finger with a varying degree of rotation and a small, dysmorphic watch-glass nail associated with swelling of the distal interphalangeal (DIP) joint. The deformity usually becomes obvious between the ages of eight and fourteen years, and twice as commonly in females. It can be progressive and sometimes is associated with swelling and redness at the base of the nail, but it is rarely painful. Kirner’s deformity is suggested to be inherited in an autosomal dominant trait with incomplete penetrance but sporadic cases were also reported. No specific genetic linkage has been identified ¹,²,³. Camptodactyly is a non-traumatic flexion contracture of the proximal interphalangeal (PIP) joints of one or more fingers, resulting in a permanent volar inclination of the affected digits. Little finger is the most frequently affected one, but additional fingers may also be involved. The degree of flexion varies from very mild to almost right-angled bend and may progress during adolescent growth period or remain unchanged for years. Camptodactyly may be sporadic or familial. In severe forms, flexion contractions are usually seen in multiple digits of both hands and this form is associated with other congenital anomalies such as chromosomal disorders, craniofacial diseases and
skeletal deformities. In isolated camptodactyly, one or both little fingers are affected and this form is inherited in autosomal dominant trait with variable penetrance or expressivity, or it may be sporadic. Knuckle pads upon the PIP joints might also be observed in combination with isolated camptodactyly.

The combination of these two entities in a family had not been previously reported. The family presented here suggests that Kirner’s deformity and camptodactyly may have a common genetic factor.

Report of the Case

A 14 year old girl presented to our clinic with a history of swelling and increasing deformity of the distal phalanx of her little fingers. The deformity has existed for 3 years. The patient had no pain but had some discomfort when typing. There was no history of trauma and infection. Physical examination confirmed palmar and radial deviation of distal phalanges of both little fingers, but the right side was the most affected one (Fig 1).

Figure 1. Radial and palmar deviation of the distal phalanx of little fingers.

The nails were also curved in volar direction. The flexor and extensor mechanisms functioned normally. Her routine blood examinations were normal and rheumatoid factor was negative. On radiographies, the head of the distal phalanges of little fingers showed a pronounced tilting in both volar and more slightly in the radial direction, with preservation of the epiphyseal, metaphyseal and articular alignment (Fig 2). This deviation was recognized as being Kirner’s deformity. The family history revealed that there were no any relatives having this deformity but there were 2 people having camptodactyly in various degrees in father’s family (Fig 3).

Figure 2. AP view (a) and lateral view (b) of right and left little fingers.

Figure 3. Pedigree of patient’s family

Figure 4. Radiograph of both hands and photograph of right and left hand of the case II.2 (A)
The patient's aunt (case II.2) had bilateral camptodactyly in her both little fingers (Fig 4) and had knuckle pads on PIP joints of all fingers except the thumbs (Fig 5). Her daughter (case III.5) had unilateral camptodactyly in only right little finger in a mild degree (Fig 6), and had knuckle pads on all fingers except thumbs. The patient said that paternal grandmother (case I.2) who was not alive and one aunt (case II.5) had also been affected but they have not been examined to confirm this.

Discussion

Kirner’s deformity and camptodactyly have been regarded as independent congenital hand deformities \(^1\)–\(^3\). They both may be sporadic or familial \(^1\)–\(^2\),\(^5\)–\(^6\),\(^8\). In inherited forms of Kirner’s deformity, it is an independent anomaly, but it can also occur as a minor part of different syndromes such as Down syndrome, pes cavus, osteomyelitis, and progressive myositis ossificans \(^2\). The etiology of Kirner’s syndrome is not well understood. Kaufmann et al. suggested a lysis between diaphysis and epiphysis of the terminal phalanx \(^10\). Several other factors as a developmental defect in enchondral ossification, an osteochondrosis possibly of vascular origin or of unknown origin \(^9\) have been suggested.

Some familial cases of Kirner’s deformity \(^1\)–\(^8\) and cases without a family history \(^7\) have been reported. Damen et al. reported a patient with Kirner’s syndrome and 7 clinodactyly cases in a family. He suggested that there could be a common genetic factor for Kirner’s deformity and clinodactyly \(^9\). There is no any report about the association of Kirner’s deformity and camptodactyly. Camptodactyly may occur as an isolated deformity as well as a part of many well-characterized hereditary syndromes. Isolated type can be sporadic or can have an autosomal dominant trait with variable penetrance and expressivity. The first locus of 5th finger of camptodactyly was mapped to 3q11.2-q13.12 in a large German family by Malik et al. and was reported that this locus was not shared by any other camptodactyly-like syndrome \(^7\). The gene for this deformity has not been identified as well as any specific gene locus and gene has not been reported for Kirner’s deformity.

We here present a family with one case of Kirner’s deformity and 2 cases with camptodactyly and 2 probable camptodactyly cases who were not examined. The presence of both camptodactyly and Kirner’s deformity in this family suggest that there could be a genetic linkage between these deformities or they could be different expressions of a common genetic disorder. As Kirner’s deformity has not been mapped to any chromosomal location, the gene responsible for this syndrome may also be located in the chromosomal locus of camptodactyly (CAMPD1). Experimental studies should be performed in families in order to define the locus and gene of Kirner’s deformity.

REFERENCES


