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Congenital contractural deformities of the fingers and arthropathy

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SUMMARY Four patients are described who presented with congenital finger contractures and arthropathy. There was synovial cell hyperplasia and giant cells but no inflammatory process. Radiographs showed flattening of the metacarpal and metatarsal heads and the proximal femoral ossification centres. In the oldest patient the process had subsided leaving slight contractures but severe impairment of hip mobility. In another the arthropathy was still prominent in the early teens. In a third, finger contractures had failed to respond to conservative or surgical measures.

Key words: camptodactyly, hypertrophic synovitis.

Congenital contractural deformities of the fingers and large joint arthropathy due to a non-inflammatory hyperplasia of synovial tissue have been described by a number of observers.1–3 We have studied two siblings and two unrelated cases with these features. In all the cases the parents were healthy, unrelated, Newfoundland Caucasians. There were no other individuals with contractures or relevant arthropathies in the three families. The purpose of this communication is to provide further observations on what appears to be a well defined syndrome.

Case reports

CASE 1
The youngest child of a family of six was the product of a normal full-term delivery. At birth her fingers were held in a clenched position; they could be passively extended but became progressively more fixed. Beginning at the age of 2 years a number of surgical attempts were made to correct the contractures in the thumbs, left ring, right index, middle, and little fingers. Hypertrophy of the sheaths, jelly-like fluid within the sheaths, and oedema or shortening, and atrophy of the tendons were recorded on different occasions in the operative descriptions. The tendon sheaths of one or more digits at a time were divided and restrictive bands removed, permitting full passive extension and tendon lengthening at two of the later operations. The results were disappointing, and on each occasion contractures returned and developed in the hitherto unaffected little fingers. In early adolescence the proximal interphalangeal (PIP) joints of the little fingers were arthrodesed.

During childhood she complained of pain in the elbows, knees, and ankles. One observer reported effusions but no heat or tenderness. At the age of 13 years she was a well proportioned child with no dysmorphic craniofacial features but with the finger deformities previously described. There was also a slight flexion contracture of the left elbow and slight contractures of the PIP joints of the toes. The range of movement in the hips was normal apart from only 10° internal and external rotation. There were no effusions. Radiographs of the hands, skull, thoracic and lumbar spine, and pelvis were unremarkable apart from flattening of the metacarpal heads and an appearance suggestive of bilateral slipped femoral epiphyses. The dermatoglyphic patterns did not show the vertical alignment of the palmar lines seen in some other types of arthrogryposis. Other laboratory investigations, including haemogram, immunoglobulins, rheumatoid factor, serum comple-
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ments urinary glycosaminoglycans, and amino acid levels, were normal or non-contributory. The fibroblast culture showed a full complement of 46XX normal chromosomes by Giemsa banding.

The tendon sheath excised at 2 years of age showed synoviocytic hyperplasia, focal stromal oedema, and surface fibrin deposition. At age 16 years the tendon sheath showed scarring. Ultrastructural examination of a tendon showed a few tendon fibrocytes, with cytoplasmic accumulation of loose filamentous strands and a mixture of densely and loosely packed fibrils. Focal disarray of collagen fibres was noted; these had a regular periodicity but were of varying diameters (50–175 nm). Amorphous electron dense finely granular material was abundant between collagen fibres, together with cellular fragments. The appearance of the skin, skeletal muscle, and fibroadipose tissue was unremarkable by both light and electron microscopy.

CASE 2
The oldest sibling of case 1 was a normal full-term baby. At birth it was noted that he did not straighten his fingers, though they could be extended passively but would return to a fixed position on release of pressure. Efforts to correct the deformity by physiotherapy and splinting were unsuccessful, and at 5½ years he was referred for surgical correction of the deformities. Swelling of the knees with patella taps and large wrists and ankles were noted. The flexor tendon sheaths were excised in digits of the left hand and a few weeks later in the thumb, index, and middle fingers of the right hand.

At the age of 23 years he began to complain of pains in the right hip which he ascribed to years of participation in contact sports. There were slight contractures of the elbows, PIP joints of the toes and the fingers, with lack of full flexion in the right ring finger. The left hip tended to rotate externally to 20° when flexed, with only a few degrees of internal and external rotation from that position. Internal and external rotation were also slightly reduced in the right hip. The knees showed a somewhat boggy synovium but were otherwise unremarkable. The peronei were thin, the interossei wasted, and the knee tendon reflexes were brisk, but otherwise the physical examination was normal.

Laboratory investigations were normal. Radiographs were normal apart from extreme lordosis at L5–S1 and an appearance in the hands and pelvis similar to that of case 1 (Figs 1 and 2).

CASE 3
She was a normal full-term delivery, the second of three siblings, the other two being healthy boys. She was born with clenched fingers and later was found to have swelling of a number of joints. Splints were prescribed to straighten the fingers with some benefit. At 6 years of age the flexor pollicis tendon sheaths were released. During childhood she complained of intermittent pain in the left leg but participated in sports.

Fig 1 AP view of hands. Flatening of the articular surface of metacarpal heads. Periarticular osteoporosis. Underdevelopment of styloid processes.
cytoplasm of some synoviocytes (but not fibrinogen and immunoglobulins A, G, and M) (Fig. 3c). Multinucleated giant cells were easily seen among synoviocytes in both superficial and deep areas (Fig. 3d). These had more or less central nuclei morphologically similar to those of adjacent synoviocytes. The cytoplasm was abundant and homogenous showing a periodic acid-Schiff reaction and the same serum components as the synoviocytes.

CASE 4
Case 4 was a full-term delivery. There were two miscarriages before the propositus and a normal male born subsequently. At the sixth week of pregnancy the mother had a slight bloody vaginal discharge and spent seven months in bed receiving medroxyprogesterone acetate every two weeks from six weeks to the eight months. At six months she began to have contractions and was then placed on Ventolin for the remaining part of the pregnancy.

The child was born with all fingers clenched. At eight months she was found to have flexion deformities of the fingers at the PIP joints and recurvatum deformity of the MCP joints. She was prescribed night splints, and by 14 months the deformities had corrected themselves except for slight contractures and crepitus of the PIP joints of the ring and little fingers. At 19 months puffiness, but no heat, of the knees and ankles was noted, as well as swelling of the extensor tendons of the left wrist and a slight contracture of the right elbow. Hip movements were normal and there were no contractures of the toes. She was placed on ASA 100 mg/kg.

Pertinent laboratory findings: erythrocyte sedimentation rate 26 mm/1st h (Westergren), IgM 185 mg% (normal 61±19), rheumatoid factor test non-reactive. Radiographs showed an increased acetabular angle with abnormal indexes suggestive of a hip dysplasia.

Discussion
The four cases presented here have the typical features of the syndrome of congenital flexion contractures of the fingers and arthropathy.1-3 They are similar to the type 1 distal arthrogryposis identified by Hall and her associates,4 though the hands of the fourth case resemble their type II E. The presence of the finger contractures at birth and the absence of inflammatory cells in the synovium distinguish them from the familial arthritis and camptodactyly reported by Malleson.5

The clinical course of the finger deformities was quite variable. In cases 3 and 4 the finger deformities responded to conservative measures, and in case 2 simple incision of flexor tendon sheaths provided

Fig. 2. AP view of pelvis at 24 years. Broadening and flattening of the proximal femoral epiphysis. The femoral necks are short and thick with a coxa vara deformity.
Fig. 3  (a) Villus showing hyperplasia of synoviocytes, with oedema and focal collagen degeneration in subsynovial zone. Inflammatory cells absent. (H and E, ×100). (b) Avascular villus with deposition of fibrin on surface and within collagenous tissues. (Immunoperoxidase staining for fibrin ×100). (c) Avascular villus showing positive cytoplasmic staining of synoviocytes and chondrometaplasia. (Immunoperoxidase staining for α, antitrypsin ×200). (d) Several multinucleated giant cells in superficial and deep synovial zones. Note central uniform nuclei and homogenous cytoplasm. (H and E, ×200).
complete correction. In contrast, conservative and surgical measures were disappointing in case 1. Congenital contractural deformities of the fingers, it is said, if left untreated tend to decrease but may be improved or worsened by corrective operations.

There are no comprehensive descriptions of the radiographic appearance of this syndrome. The distinctive features in our cases were flattening of the proximal femoral ossification centres similar to that reported by Jacobs, together with flattening of the metacarpal and metatarsal heads, which was not present in Athreya’s patients but has been reported in the Morquio syndrome.

The syndrome of congenital flexion contractures of the fingers and arthropathy is said to affect larger joints and to subside early in infancy leaving only residual contractures and restricted movements in the hips. Three of our cases (1–3) were followed up long enough for an opinion to be formed on the course of the arthropathy. Two of them followed the expected course, but in the third (case 3) a florid polyarthritis affecting both large and small joints was still present in her early teens. Thus the arthropathy may be variable.

The syndrome of congenital camptodactyly and arthropathy begins in utero and affects the synovium of joints and tendon sheaths. The distinctive pathological features are synovial cell hypertrophy and hyperplasia, multinucleated giant cells, avascular villi, and absence of inflammatory cells. These changes were noted in those of our patients where histological material was examined. Athreya noted fibrin deposition on some of the villus surfaces, and we were able to confirm this finding (Fig. 3b). We also observed α₁ antitrypsin (a marker for histiocytes) in both single nucleated and multinucleated giant cells (Fig. 3c).

The finger contractures of the congenital camptodactyly and arthropathy syndrome are considered to be the result of a tenosynovitis causing disruption of tendons and subsequent scarring and adhesions between tendons and tenosynovia. At various times different phases of this sequence were observed in our first case. Electronmicrographs of a tendon showed collagen fibres of regular periodicity but variable diameter. Some of the tendon cells had the appearance of myofibrocytes, suggesting that they are metabolically active and similar to the myofibroblasts seen in reparative granulation tissues. These ultrastructural findings, taken in conjunction with Athreya’s observation of intracellular collagen fibres, could be interpreted as pointing to a defect in collagen synthesis as a component of the syndrome.

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