Arthrogryposis, ophthalmoplegia, and retinopathy: confirmation of a new type of arthrogryposis

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Abstract
Arthrogryposis multiplex congenita is a heterogeneous condition and many different types are clinically recognisable. Recently, a new type of autosomal dominant arthrogryposis was described in a father and son. We report on a male patient with similar clinical features, confirming this distinct type of arthrogryposis. The condition is characterised by congenital contractures of the hands and feet with diminished or absent phalangeal creases, ophthalmoplegia, a rigid trunk, deep set eyes, and (in the oldest patient) an abnormal electroretinogram. Differential diagnosis from amyoplasia, the different types of distal arthrogryposis, and symphalangism is discussed. (J Med Genet 1993;30:78-80)

Arthrogryposis multiplex congenita has been defined as "congenital, non-progressive limitation of movement in two or more joints in different body areas".1 Over the past 10 years, many different clinical types of arthrogryposis have been delineated2-4 and recently an apparently new type of arthrogryposis was reported in a father and son.5 We had the opportunity to observe the same combination of clinical signs and symptoms in a male patient, confirming the existence of this new entity.

Case report
The proband is the second son of healthy, non-consanguineous Dutch parents. His older brother is healthy. Maternal age at the time of birth was 27 years and paternal age 35 years. A cousin (the mother's sister's son) was born with an isolated right club foot. Pregnancy was complicated by hyperemesis gravidarum but no medication was taken. Intrauterine position was constant. The mother did not recall specific details of fetal movement or the amount of amniotic fluid. The birth was normal (exact position not known) at 36 weeks of gestation, birth weight 2700 g. At birth rigid fingers and bilateral club feet were noticed. In the neonatal period a hypertrophic pylorus was surgically treated. Several orthopaedic operations were performed on both feet, the last time being a triple arthrodesis on both feet in 1982. Despite the medical problems psychomotor develop-
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Figure 3  Face and profile of the patient showing the deep set eyes and broad mouth with thin lips.

ment was normal. A family photograph at the age of 8 months showed the deep set eyes and rigid fingers (fig 1). At the age of 6 years he had a triangular face, deep set eyes, and prominent ears. At the age of 17 years the proband was referred for genetic counselling with a diagnosis of 'arthrogryposis multiplex congenita'. On clinical evaluation (fig 2) he was a lean, rigid boy with dry skin. He walked stiffly. Height was 170 cm (3rd to 10th centile), weight 48 kg (3rd to 10th centile for height), and head circumference was 55 cm (10th centile). Both eyes were deep set and there was ptosis (fig 3). The palpebral fissures measured 2.6 cm (normal) and inner canthal distance 3.1 cm (normal). There was little facial expression although motility was normal. He was unable to move his eyes laterally or to look upwards. Muscle mass was reduced especially in the lower limbs; the shoulders were hunched and antverted and the back was rigid. He had pectus excavatum. The fingers were long and the phalangeal creases were totally absent. Flexion was limited to about 30° (fig 4). Both lower legs were thin, the left side more pronounced than the right. The feet showed multiple scars from the orthopaedic surgery. Apart from the ophthalmoplegia, no further neurological abnormalities were found. Intelligence was normal. Ophthalmological investigation showed small eyes with ambylopia in the left eye and 6/10 vision in the right eye. A central scotoma could be seen in both eyes and abnormal pigmentation was present in both retinal maculae. Both eyes showed a Duane anomaly. On electrophysiological examination, the pattern reversal visual evoked response showed less response than normal, but was otherwise unremarkable. The flash visual evoked response was also normal. The electroretinogram failed to give any result because of practical problems (painful test for the patient because of the small, deep set eyes). Chromosomes were normal male 46,XY. Creatine kinase levels were normal. Electromyography showed no neurogenic or myopathic abnormalities and conduction velocities of the nerves were normal. Radiological examination of the vertebral column showed normal vertebrae with limitation of flexion. Radiology of the hands and wrists showed normal carpal bones without fusion, narrow interphalangeal joints, and no bony symphalangism (fig 5). Hearing was normal. CT scan of the brain and orbits showed no abnormalities of the brain and normal eye muscles. CT scan of the muscles of the upper arm, lower arm, thigh, and lower leg showed no abnormalities.

Discussion

The clinical features of this patient are very similar to those in the father and son reported by Lai et al.7 (table). The facial resemblance of the son and the present patient at the age of 6 years is striking.

For the differential diagnosis, amyoplasia, distal arthrogryposis, and symphalangism were considered in the present patient. The reduced muscle mass, the hunched, antverted shoulders, the stiffness of the spine, and the club feet were very suggestive of amyoplasia; this diagnosis was initially made in the father described by Lai et al.7 The ophthalmoplegia, deep set eyes, ptosis, and retinopathy, however, are incompatible with this diagnosis.4 In the distal arthrogryposes, a type I and type II have been delineated.3 In type I no associated anomalies are present, intelligence is normal, and inheritance is autosomal dominant. In type II subdivisions have been made (type II A-E) according to the different associated findings. Presumed new forms of type II distal arthrogryposis have since been reported but we could not fit the clinical data of our patient into any of the clinical types of distal arthrogryposis.6

Symphalangism can be defined as bony or fibrous ankylosis of interphalangeal joints resulting in a rigid digit.8 In our opinion, bony fusion is mandatory for the diagnosis of symphalangism in an isolated patient. In the present patient the interphalangeal joints were narrow but the phalanges were not fused, thus
on clinical grounds the diagnosis of symphalangism was excluded.

Different eye abnormalities have been described in patients with arthrogryposis multiplex congenita but we could find no patients who combine the eye anomalies and joint limitations as seen in the three patients with the type of arthrogryposis described here and by Lai et al. A mitochondrial myopathy was suggested in the patients reported by Lai et al because of the combination of (possibly progressive) ophthalmoplegia and muscular abnormality. In the present patient, followed up for three years, we have no indication of progression of the condition. In our opinion a mitochondrial disturbance is not likely in our patient.

Inheritance in the father and son described by Lai et al undoubtedly is autosomal dominant. Family history in the present patient was not informative although a maternal cousin had serious unilateral club foot. No clinical signs of the condition were found in the patient and the healthy brother. Paternal age at the time of birth was 35 years, suggesting a new dominant mutation in the patient.

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