A three generation family with fibrodysplasia ossificans progressiva

J M Connor, H Skirton, P W Lunt

Abstract
A family with five persons affected with fibrodysplasia ossificans progressiva (myositis ossificans progressiva) in three generations is described. This is the first well documented three generation family with this condition and provides further evidence for autosomal dominant inheritance. A wide range of phenotypic severity is apparent, from disabling ectopic bone formation and premature death to an asymptomatic adult with characteristic big toe malformations. (J Med Genet 1993;30:687-9)

Fibrodysplasia ossificans progressiva (FOP, myositis ossificans progressiva, MIM 135100) is a rare disorder in which physical handicap owing to progressive soft tissue ossification accompanies characteristic skeletal malformations. Over 600 patients have been described and attempted total ascertainment in the UK found a minimum point prevalence of one patient per 1·64 million of the population. The sex ratio is equal and autosomal dominant inheritance is favoured on the basis of concordant monozygotic twins, a paternal age effect for presumed new mutations, and several instances of parent to child transmission, including male to male transmission. Only one three generation family with FOP has been published. Gaster in 1905 mentioned a family with a grandfather, father, and three sons affected in a discussion at a medical meeting but never published the family in further detail. We thus wish to describe a contemporary three generation family with FOP.

Case report
The family pedigree is shown in fig 1. The proband (II.2) was asymptomatic until 22 years of age when she developed jaw fixation after extraction of wisdom teeth. She had a further operation at 23 years of age to replace the right temporomandibular joint with a Bowerman-Conroy titanium prosthesis but jaw fixation recurred one week afterwards. She is now 47 years of age and her only symptom has been gradually increasing back stiffness in recent years. On examination she has limited neck mobility and jaw fixation but normal mobility at other joints and clinically normal big toes. A radiograph of her cervical spine showed anterior osteophytic bridging of several vertebral bodies (fig 2) and a radiograph of her feet showed symmetrical bony spurs on the medial sides of the heads of the first metatarsals (fig 3).

Her father (I.1, fig 1) had also been asymptomatic until he developed back and neck stiffness at 2 years of age when he had an anterior osteophyte of the cervical spine removed. He had further operations at 13 years of age for anterior osteophyte removal of the first cervical vertebra and then at 26 years of age for a failed fusion at the atlanto-axial level. He is now asymptomatic.

Her grandfather (I.1, fig 1) had no symptoms and died at 60 of cancer. He had been assessed at the age of 26 years with a physical examination but no radiological assessment. She had a brother (III.1, fig 1) who is now 20 years of age and asymptomatic.

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stiffness after trauma. His jaw also became fixed after trauma and by his forties he had developed a limp. He remained ambulant (with the use of a stick) until his early seventies and died at 72 years of age from a myocardial infarction. The diagnosis of FOP was confirmed at the Royal National Orthopaedic Hospital, London and some of his x rays are still held in their radiological museum. These were reviewed and showed normal hands, malformed big toes with superimposed ankylosis (similar to his granddaughter III.1), progressive ankylosis of the cervical spine, and multiple areas of soft tissue ossification.

The proband’s sister also had FOP. She was well until 15 years of age when spontaneous swelling of the left leg occurred. A biopsy was performed and the family was told she had a fibrosarcoma. The swelling subsequently subsided but she had limited left knee movements thereafter. In her late teens she developed limitation of movement at the right elbow after trauma. In her twenties she developed a succession of painless lumps on her back. After this she started to walk with a frame. She gradually became more disabled and was bed bound for a year before her death from pneumonia at 28 years of age. Her case records and radiographs are no longer available but her hospital diagnosis at the time of death was myositis ossificans.

The proband’s two daughters also have FOP. The older (III.1) has had stiff big toes for many years but is otherwise asymptomatic. Her examination at 24 years of age was normal except for no mobility at the interphalangeal joints of both big toes. Her radiographs showed malformed halluces with bony spurs on the medial sides of the heads of the first metatarsals and ankylosis of the halucal interphalangeal joints (fig 4). Her radiograph of the cervical spine was normal (fig 5). The younger daughter was well until 13 years of age when she first noted painful lumps on her back. These appeared over several hours and resolved over several days. Subsequently she has noted stiffness of her neck and lower back. For the past three years she has experienced painful limitation of movement of both hips but especially the right. She noted pain and clicking when she walks and this had been associated with occasional painful locking. On examination at 23 years of age she had palpable ectopic bone in the left lumbar region, limited mobility of the neck and spine, and painful limitation of all movements at both hips (right more than left), but clinically normal big toes. Her radiographs showed malformed big toes with medial spurs on the heads of the first metatarsals (fig 6), an ectopic bony bar in the left lumbar region (fig 7), a normal cervical spine (fig 8), and abnormalities of both hips with multiple osteochondromata and deformed femoral heads (fig 9).

Discussion
This family shows many typical features of FOP but has two unusual features and one
Hall and Sutcliffe\textsuperscript{11} described abnormal cervical vertebrae with small bodies, enlarged pedicles and short 'massive' spinous processes in eight children with FOP. Similar changes were found in four of eight other children with FOP and in adult life variable fusion of the cervical vertebrae was a consistent feature.\textsuperscript{12} This fusion was first noted between adjacent neural arches in late childhood but also involved the vertebral bodies in some adult patients. The normal cervical spines in adulthood in patients III.1 and III.2 are thus unusual and probably reflect the generally benign clinical course in these patients.

In general patients are severely handicapped owing to ectopic bone formation. Most patients have severe limitation of the spine and shoulders by 10 years of age, one or both hips are involved by 20 years of age, and most patients are chair or bed bound by 30 years of age.\textsuperscript{10} The natural history in II.3 is thus fairly typical whereas the four other affected subjects in this family have had considerably milder clinical courses. This may reflect the nature of the underlying mutation or factors affecting variable expression of this dominant trait. As yet these factors are unknown and no medical treatment has been shown to influence the long term prognosis.\textsuperscript{1}

Multiple osteochondromata have not previously been described in the hip joint in FOP. Usually these represent metaphysic cartilaginous nodules derived from the synovial membrane (synovial osteochondromatosis).\textsuperscript{13} Synovial osteochondromatosis most commonly occurs in young or middle aged adults and the most frequent areas of involvement are the knee joint, the hip, elbow, and shoulder.

The genetic defect causing FOP and its chromosomal regional localisation are currently unknown and linkage studies in families such as this will assist with exclusion mapping of possible candidate genes for this disorder.

We wish to thank Dr D J Stoker of the Royal National Orthopaedic Hospital Trust for kindly providing us with copies of radiographs of I.1.

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