Rapidly destructive hip disease following ipsilateral hemiparesis: report of two cases

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SUMMARY Two patients who developed rapidly destructive arthropathy of the hip following ipsilateral hemiparesis are described. The possible significance of this association is discussed.

Key words: osteoarthritis, coxarthrosis, analgesic hip, destructive arthropathy, apatite associated destructive arthropathy, Charcot arthropathy.

There have been several reports concerning the effects of neurological deficit on the subsequent development of a variety of joint diseases.1-10 Most commonly it has been concluded that either a central1-5 or peripheral6-10 lesion protects the affected limb from subsequent development of arthropathy. This sparing effect has been attributed to disuse with subsequent reduction in damaging mechanical factors.2 3 10 The converse effects of neurological deficit on established joint disease are less well recorded; there are two reports, however, of hemiparesis resulting in a flare of rheumatoid synovitis confined to the paretic side.11 12

Apatite associated destructive arthropathy is a recently recognised form of 'joint failure' characterised by severely painful and rapid progression to joint destruction. Radiographic features include marked cartilage and bone attrition with paucity of osteophytes and cysts.13 Synovial fluid from such joints is typically non-inflammatory and contains plentiful alizarin red staining particles identified as apatite.13 14 The predilection for this condition to affect large joints of the elderly is striking,13-16 but the pathogenesis remains obscure.13 14

Two elderly patients who developed unilateral rapidly destructive coxarthrosis following ipsilateral hemiparesis are described. The clinical, radiographic, and histological characteristics of their arthropathy were typical of those described for apatite associated destructive arthropathy. The relevance of this observation to the pathogenesis of rapidly destructive arthropathy is discussed.

Case reports

CASE 1
A previously fit 58 year old right handed window cleaner developed an acute left hemiparesis. He made an excellent recovery but was left with mild residual hemiparesis and increased tone on the left side. Hypertension was diagnosed and adequately treated by bendrofluazide 5 mg daily. Fifteen years later (age 73 years) he developed insidious onset of pain and stiffness in the left hip; radiographs showed bilateral superior pole joint space narrowing with sclerosis and osteophyte formation. His symptoms were readily controlled by physiotherapy and naproxen 500 mg twice a day.

At age 77 years he developed a second, more severe hemiparesis causing profound weakness and transient sensory disturbance, again on the left side. He made only slow improvement during the next three months and complained of severe exacerbation of left hip symptoms on attempted weight bearing, and developed night pain sufficient to interrupt sleep. His symptoms did not respond to naproxen 1 g daily and paracetamol 4 g daily, and in an attempt to control his severe pain he inadvertently took multiple drug therapy: for four weeks he took naproxen 1 g and indomethacin slow release 150 mg daily. Eight months after onset of this severe pain he was admitted to hospital with bleeding from multiple superficial gastric erosions. There was no history of significant trauma nor had he received steroids. Examination at this time showed marked painful reduction in all movements of the left hip. There was mild, pain free restriction of internal rotation and flexion of the right hip, but apart from several Heberden’s nodes, more marked on the
right, there was no evidence of arthropathy elsewhere. There was 4/5 pyramidal weakness affecting the left arm and leg, but no sensory abnormalities. Radiographs showed the previously noted changes of superior pole osteoarthritis on the right; the left, however, showed apparent widening of the joint space, marked bone attrition affecting both femoral and acetabular components, and absence of osteophytes and cysts (Fig. 1). Plain radiographs of knees and hands (to include wrists) showed mild changes of osteoarthritis in several interphalangeal joints: no chondrocalcinosis was apparent. Investigations included; haemoglobin 12·8 g/dl (128 g/l), erythrocyte sedimentation rate 37 mm/h, serum urea 11·1 mmol/l (normal range 1·0–6·5 mmol/l), serum creatinine 221 μmol/l (normal range 60–120 μmol/l), serum electrolytes, glucose, magnesium, calcium, uric acid, iron, iron binding capacity, ferritin, vitamin B₁₂, red cell folate, liver and thyroid function tests were within normal limits. Rheumatoid factor and serological tests for treponemal infection were negative.

Sequential use of various non-steroidal anti-inflammatory agents and analgesics failed to control his symptoms, and on general medical grounds he was considered unsuitable for hip replacement.

Over the succeeding nine months, however, his hip pain steadily improved, despite further radiographic progression (Fig. 2). Twelve months after the second hemiparesis he is ambulant and independent, experiencing hip pain only after prolonged weight bearing.

CASE 2

A 68 year old right handed woman developed an acute onset of left hemiparesis. She had received bendrofluazide 5 mg daily for hypertension for three years, and had a three year history of bilateral hip pain managed successfully by weight reduction, physiotherapy, and occasional paracetamol. Hip radiographs performed two years before her hemiparesis showed moderate bilateral osteoarthritis with central narrowing, sclerosis, and cyst formation. She made good functional recovery from her hemiparesis but was left with residual increased tone, brisk reflexes, extensor plantar response, and mild spastic gait on the left. No sensory signs were elicited at any stage, and the pyramidal weakness had largely resolved after two months.

Three months after the onset of her hemiparesis she developed marked increase in left hip symp-
toms, with predominant pain on usage and moderate night pain. There was no obvious predisposing trauma. Attempts to control her severe progressive pain with non-steroidal anti-inflammatory agents and analgesics were unsuccessful, and she was referred for orthopaedic opinion. Examination three months after exacerbation of symptoms disclosed marked painful restriction of all left hip movements. The right hip showed only pain free limitation of internal rotation and flexion and apart from mild bilateral Heberden’s nodes there was no evidence of arthropathy elsewhere. No new neurological signs were elicited. Hip radiographs showed no change from the previously recorded osteoarthritis on the right, but on the left there was severe destructive change affecting both femoral and acetabular components (Fig. 3). Haematological and biochemical screening showed no underlying predisposing abnormality.

At total hip replacement five weeks later marked attrition of cartilage and bone was confirmed macroscopically. Histological examination showed hyperplastic villous synovial tissue containing chronic inflammatory cells, macrophages laden with haemosiderin (Perl’s stain), and occasional multinucleate giant cells. Numerous irregular intrasynovial bony fragments were present within collagen staining material, the latter often hyalinised. Alizarin red staining showed further, smaller calcific particles, which frequently coalesced with van Gieson’s stain. No birefringent crystals were identified and there was no microscopic or culture evidence of tuberculous or pyogenic infection.

Nine months after surgery the patient remains mobile, independent, and pain free.

**Discussion**

Two patients who developed rapidly destructive hip disease following ipsilateral hemiparesis are described. Both had recorded pre-existing hip osteoarthritis which had been stable before the onset of the neurological event. Neither patient had received steroids or sustained previous hip trauma, and a full clinical and metabolic screen failed to disclose associated or underlying disease. In the absence of other recognised predisposition to rapidly destructive arthropathy, the close temporal relationship to hemiparesis, the previously quiescent joint disease, and the striking lateralisation strongly support a causal rather than chance association. To our knowledge this is the first report of such an occurrence.

A number of reports describe rapidly destructive arthropathy of large joints under a variety of titles including ‘analgesic hip’, ‘indomethacin hip’,17–22 apatite associated destructive arthropathy,13 Milwaukee shoulder syndrome,14 lytic arthropathy,23 senile haemorrhagic arthrosis,24 and idiopathic destructive arthropathy.25 The presence of plentiful apatite crystals in synovial fluid and tissues has been emphasised,13 14 26 but the pathogenic significance of these particles in rapid joint destruction remains unknown.13 14 26 27 Although the factors that predispose to rapid joint destruction are unclear, advanced age, being female, and the occurrence of calcium pyrophosphate dihydrate crystal deposition either within the involved joints or at distant sites have all been incriminated27–30; the possible contributory role of non-steroidal anti-inflammatory agents remains controversial.31 Differential diagnoses that may be considered include late avascular necrosis and sepsis, though the clinical, synovial fluid, and radiographic features most closely resemble Charcot arthropathy.31 Two radiographic appearances of Charcot arthropathy are recognised; a ‘hypertrrophic’ type characterised by exuberant sclerosis, osteophyte formation, abundant osseous debris, and dissolution of normal joint architecture; and an ‘atrophic’ form with extensive bone resorption without evidence of accompanying bone repair.32–35 It has been suggested that the atrophic form represents an earlier stage of the disease.

**Fig. 3** Left hip radiograph of patient 2 taken six months after onset of hemiparesis, showing severe atrophic disease with marked attrition of both acetabular and femoral components.
before repair has begun. There are two main theories of causation: firstly, that reduced pain and proprioception lead to excessive mechanical trauma that disrupts joint tissue, and secondly, that abnormality of neurovascular control disturbs bone and cartilage nutrition. The importance of adjacent bone fracture, a not uncommon early feature, remains uncertain. Although generally regarded as a characteristically pain free condition, pain may be a prominent symptom in up to 30% of patients and was indeed emphasised by Charcot in his original description.

In our two patients their advanced age, the rapid severe clinical and radiographic progression, the individual atrophic x ray features, the synovial histology in case 2, and the absence of sepsis or other cause of joint destruction are fully consistent with a diagnosis of apatite associated destructive arthropathy, idiopathic destructive arthropathy, or 'analgesic hip'. The radiographic appearance in isolation is also indistinguishable from atrophic Charcot arthropathy: clinical points of differentiation, however, were the prominence of pain and the restriction rather than increase of joint mobility. Radiographic similarity between apatite associated destructive arthropathy and atrophic Charcot arthropathy has previously been noted, and it is therefore of interest that apatite associated destructive arthropathy, an uncommon arthropathy of the elderly, should develop in the hips of both our patients following ipsilateral hemiparesis. Rapidly destructive arthropathy is not a recognised complication of hemiparesis: both our patients had predominant motor impairment, only transient clinical sensory abnormality, and severe pain which inhibited even normal loading of the joint. A damaging effect from the non-steroidal anti-inflammatory agents used by these patients cannot be excluded, though the marked asymmetry coinciding with the neurologically affected side suggests a more important neurogenic cause.

Although previous reports emphasise a predominantly sparing effect of neurologic deficit on subsequent development of new rheumatic disease, the effect of hemiparesis on pre-existing joint disease may be to cause exacerbation. Both our patients had bilateral osteoarthritis of the hip and shortly following hemiparesis the hip on the affected side entered a rapidly destructive phase with marked bone and cartilage attrition. Although we cannot deduce the precise relation between hemiparesis and rapidly destructive arthropathy in these two cases, the abrupt onset, rapid progression, severe tissue loss, and non-inflammatory features are most compatible with a neurovascular theory of causation. More subtle age related neurovascular abnormality may well underlie the pathogenesis of apparently idiopathic destructive arthropathy of the elderly.

The marked clinical improvement, despite radiographic progression in patient 1 (Figs 1 and 2), is of further interest and contrasts with the poor outcome reported in patients with apatite associated destructive arthropathy. The second patient underwent surgery shortly after entering the rapid phase of joint destruction. Long term prospective studies are still required to determine the natural history of idiopathic destructive arthropathy of the elderly.

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