Trauma and seronegative spondyloarthropathy: rapid joint destruction in peripheral arthritis triggered by physical injury

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SUMMARY Two B27 positive patients developed peripheral arthritis immediately after a significant musculoskeletal injury. Unlike previously reported peripheral arthritis precipitated by trauma in B27 positive subjects the arthritis was rapidly destructive.

Key words: ankylosing spondylitis, trauma, HLA-B27 antigen.

The role of trauma in the initiation of seronegative spondyloarthropathy has been considered in various reports in the last few years.1-4 Although physical injury does not appear to determine spinal involvement in ankylosing spondylitis but brings it to the patient’s attention, probably because of the consequent immobilisation,1 there is evidence that trauma may influence the onset in B27 positive subjects of a peripheral arthritis predominantly involving the injured joints.2-4

The present paper describes two more cases of peripheral arthritis triggered by trauma in B27 positive subjects. Unlike previous reports the arthritis was rapidly destructive.

Case reports

CASE 1
A 13 year old girl was well until 26 April 1986 when she fell and struck her right hip while running in an obstacle race. X rays of the hips were normal except for a sclerosis of both acetabular roofs, which was ascribed to scoliosis (Fig. 1a).

The following day the patient had a sore throat and a high temperature. Three days later she developed arthritis of the left elbow, the right hip, and both ankles and was admitted to hospital.

Investigations showed an erythrocyte sedimentation rate (ESR) (Westergren) of 112 mm/1st h. C reactive protein (CRP) 95 mg/l (normal <5 mg/l), white cell count 11·2x10⁹/l with 70% polymorphonuclear leucocytes. The antistreptolysin O titre was

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Fig. 1a
A total dose of 800 mg of prednisone was received by the patient.

In September 1986 because of recurrence of pain in the right hip and progressive limitation of joint mobility the patient was referred to us.

Physical examination showed tenderness and severe limitation in the range of movement of the right hip and pain on movement of the left elbow.

Laboratory examination showed an ESR of 20 mm/1st h, a CRP of 14 mg/l, and negative tests for antinuclear antibodies and rheumatoid factor. HLA typing was positive for the B27 antigen.

The past medical and family history were negative for HLA-B27 associated diseases.

Roentgenography showed severe concentric loss of the joint space and erosions of the right hip and a reduction of the joint space of the left elbow. The lumbar spine and sacroiliac joints were normal. Tomography showed more clearly erosions of both the acetabular roof and the femoral head (Fig. 1b).

CASE 2
In August 1986 a 25 year old woman was hit by a falling bookcase while she was writing in her office. She sustained multiple contusions. x Rays of her right hip and femur taken the same day were normal (Fig. 2a).
During the following two months the pain became gradually more severe and disabling, so that the patient could not walk without a crutch. At the beginning of November 1986 she attended our unit.

Her past history showed that she had had three episodes of sciatic pain in the last four years and that she had felt continuous inflammatory low back pain and stiffness over the last year. There was no history of diarrhoea, urethritis, eye symptoms, or mucocutaneous lesions. The family history was negative for HLA-B27 associated diseases.

Physical examination of her right hip and lumbar spine was not possible because of the severity of hip pain. Chest expansion and cervical spine mobility were normal.

Laboratory evaluation showed an ESR of 45 mm/1st h and a CRP of 35 mg/l. HLA typing was positive for the B27 antigen.

Roentgenography showed bilateral sacroiliitis, already present in the radiogram obtained on the day of the trauma two months before, and a concentric reduction of the joint space and erosions of the right hip (Fig. 2b). The lumbar, cervical, and dorsal spine films were normal.

**Discussion**

Both these patients have a seronegative spondyloarthropathy. The first patient has a juvenile onset HLA-B27 associated asymmetric oligoarthritis. She lacked rheumatoid factor and antinuclear antibodies. The second patient has a bilateral sacroiliitis. She is B27 positive and has a four year history of sciatic pain and inflammatory low back pain and stiffness.

There seems little doubt about the close relation between the onset of peripheral arthritis and physical injury in these two patients. The first patient had arthritis of the left elbow, the right hip, and both ankles three days after trauma to the right hip. The second developed arthritis of the right hip immediately after an injury mainly involving the same joint.

The role of trauma in the onset of peripheral arthritis in seronegative spondyloarthropathy has been discussed in various recent reports. Wisniewsky in 1984 reported the cases of two B27 positive patients who developed peripheral arthritis with features of Reiter’s syndrome shortly after physical injury. Masson et al in 1985 described three more cases of Reiter’s syndrome precipitated by trauma. All three patients were B27 positive and all three developed sacroiliitis in the months and years following the trauma. In their review of 58 cases of juvenile onset B27 positive spondyloarthropathy Jacobs et al found five children who had experienced trauma sufficiently severe to consult a physician before the onset of peripheral arthritis. Other cases of Reiter’s syndrome influenced by trauma had previously been reported by Doury et al, Bernard et al, and Noer. HLA typing was not carried out in these cases.

It has been claimed that trauma also influences the onset of spinal involvement in ankylosing spondylitis. A recent study of Jacoby et al suggests that injury does not initiate ankylosing spondylitis but brings it to the patient’s attention, probably due to enforced immobilisation.

Unlike previously reported cases of peripheral arthritis triggered by trauma in B27 positive subjects, our two patients showed a rapid destructive process. Both patients showed concentric joint space narrowing and erosions in the injured joint only a few months after the trauma. The first patient also developed articular cartilage loss in the left elbow. The radiological appearance of the right hips
in both patients corresponds to those of hip involvement in ankylosing spondylitis.13 14

Cases of post-traumatic chronic arthritis of the knee in psoriatic patients have been observed by Wright,15 and the case of a patient with psoriasis and post-traumatic erosive arthritis mainly of the injured joints has also been reported by Williams and Scott.16 HLA typing was not carried out in these cases.

The mechanisms involved in peripheral arthritis precipitated by trauma in patients affected by seronegative spondyloarthritis are not clear. Wisnesky postulated that self antigens exposed after injury may lead in predisposed individuals to the development of an autoimmune reaction.2

References

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