ADVERSE DRUG REACTION

Indomethacin induced avascular necrosis of head of femur

K R Prathapkumar, I Smith, G A Attara

Abstract
Chemically induced avascular necrosis of bone is a well documented entity. Indomethacin is one of the causes of this condition but is often difficult to recognise. Review of the literature shows that only one case of indomethacin induced avascular necrosis has been reported in the English language between 1966 and the present.

The case of a young healthy man, who developed avascular necrosis of head of femur after prolonged administration of indomethacin, is reported here.

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Case report
A 33 year old postman presented with pain in his right hip. He had previously suffered from sciatica secondary to an L4/L5 intervertebral disc prolapse. He took indomethacin 150 mg daily for eight months before having laminectomy and discectomy. After surgery he was free of sciatic pain.

Four months after surgery he developed pain in the right hip. Radiography of the right hip taken at that time was normal. Indomethacin was again prescribed for two months with no relief of pain. He was not on steroids or any other medication and did not have any trauma to his right hip. Six months later he attended the orthopaedic clinic and at this stage clinical examination revealed a 15° fixed flexion deformity of the affected hip and real shortening of 1 cm. All movements of the right hip were restricted by pain.

All other joints were normal as was the general systemic examination. Radiography (fig 1) of the hip revealed collapse of the right femoral head due to avascular necrosis. His haematological and biochemical tests were normal. Magnetic resonance imaging (MRI) (fig 2) of his right hip revealed evidence of avascular necrosis while that of the left hip was normal.

Discussion
There are several causes of avascular necrosis of bone. But many cases are classified as idiopathic because of the difficulty in determining the cause. Chemically induced avascular necrosis of bone is well documented and indomethacin is one of the analgesic anti-inflammatory agents implicated (box 1). The occurrence was first documented in 1968.¹ The rate of articular destruction in indomethacin induced arthropathy is slower than that in steroid induced arthropathy.² In this case, the patient took 45 g of indomethacin for a period of 10 months, before radiological diagnosis of avascular necrosis of head of femur was confirmed. This compares with the first reported case of avascular necrosis of head of femur, after indomethacin intake, in which a total amount of 105 g was implicated as the cause of the disease.¹

It has been presumed that the effect of indomethacin is primarily brought about by its analgesic effect, which predisposes to microfractures of weight bearing joints. Inhibition of healing of these microfractures may be important in pathogenesis of indomethacin induced avascular necrosis. Indomethacin reduces sulphate uptake in the articular cartilage of experimental animals.³ The sulphate uptake of chondrocytes reflected their protein-polysaccharide synthesis, an important constituent of cartilage matrix. Indomethacin has
also been shown to produce a statistically significant depression of sulphated glycosaminoglycan synthesis in cartilage of the femoral head. These sulphated glycosaminoglycans and protein-polysaccharide compounds are important in the production of woven fibroid bone, which is the main factor in the fracture healing. Therefore indomethacin has the dual effect of predisposing the weight bearing joints to microfractures and concurrently inhibiting the healing of these fractures.

Solomon’s criteria (box 2) are an aid to making the diagnosis of drug induced avascular necrosis of bone and patients on long term indomethacin should be followed up regularly for early detection of avascular necrosis in weight bearing joints.

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