

The Torg-Winchester form of hereditary osteolysis: Orthopaedic manifestations and management

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Abstract

The genetic osteolyses are an uncommon group of skeletal disorders in which severe orthopaedic complications can occur. In this article we present the clinical and radiological features of a boy with the Torg-Winchester form of osteolysis and discuss the orthopaedic management of his upper and lower limb deformities. Heritable osteolysis warrants consideration as a possible diagnosis in children presenting with clinical deformity following minimal trauma and radiological evidence of progressive dissolution of bone.

Key words:

genetics, orthopaedics, osteolysis, skeletal syndromic

Introduction

The Hereditary Osteolyses are a group of rare genetic disorders characterised by dissolution of the bones and joints.¹ As these conditions are progressive and cause considerable physical handicap, ongoing orthopaedic measures are crucial^{2,3} for effective management.

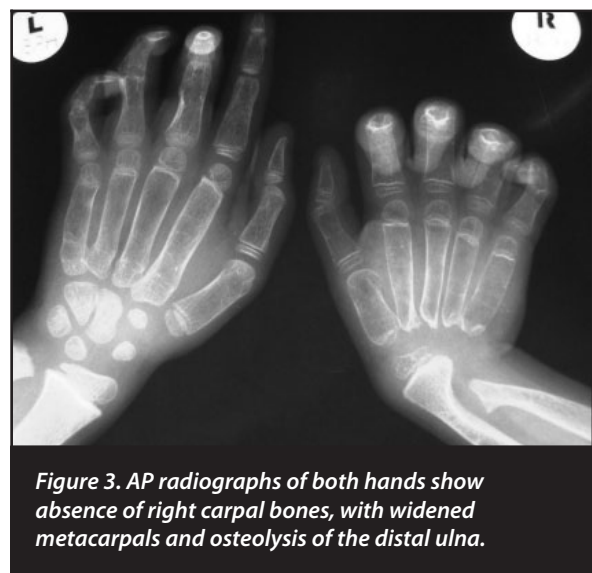
In the Torg-Winchester form of osteolysis (MIM 259600) the wrists and ankles are mainly affected, with eventual disappearance of the bones of the carpus and tarsus.⁴ The Nodulosis-Arthropathy-Osteolysis (NAO) syndrome has been incorporated into this disorder, which is now regulated as a spectrum caused by different mutations within the matrix metalloproteinase gene *MMP2*.⁵⁻⁷

In view of the rarity of this condition and the paucity of literature pertaining to the orthopaedic management of the osteolyses, we have presented, depicted and discussed our approach to this boy's skeletal problems.

Case report

A Zulu boy aged 6 years presented with complaints of a painful right leg and arm with associated deformity. His development was normal up to the age of 3 years, when a deformity of his right elbow was noticed two months after a fall. At 6 years of age he developed deformities of both hands, as well as the right knee following minimal trauma. He had no family history of similar abnormalities.

He was a healthy looking and intelligent boy, and his weight and height were normal for his age. His spine was normal, and he had no subcutaneous nodules. His musculoskeletal pathology was predominantly right-sided, but the hands and feet were affected bilaterally (*Figure 1*). He was unable to walk without assistance, due to flexion deformities of the right knee and elbow.



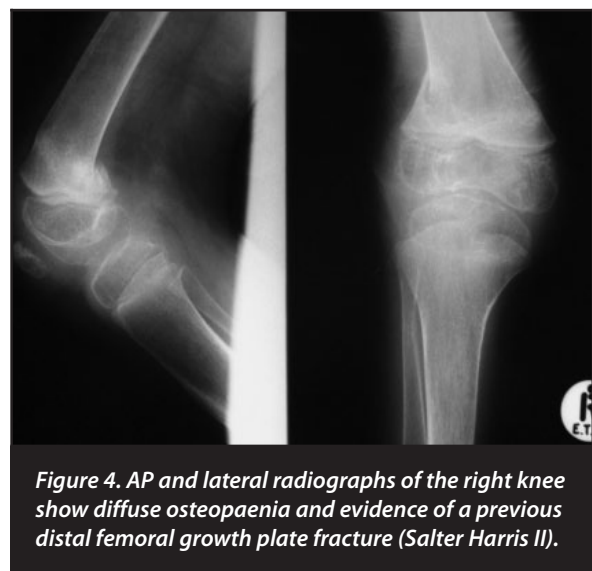
Radiographs revealed generalised skeletal lucency with absence of the carpal bones in the right wrist and destruction of the right elbow joint. The metacarpals, metatarsals and phalanges of both hands and feet were widened, with loss of tubulation. The medial regions of both clavicles were expanded. The pertinent clinical and radiological findings are depicted in *Figures 1–6* and listed in *Table I*.

Routine haematological and biochemical investigations yielded normal results and there was no biochemical evidence of parathyroid dysfunction. His renal function was normal. Facilities for the investigation of the causative *MMP2* gene were unavailable.

Orthopaedic management

Management of his upper limb pathology was focused on his right forearm and both hands. We did not consider interfering with his right elbow deformity, as the position would allow him to use his hand to feed himself. Management of the ulna club hand involved a distal ulna osteotomy and wrist spanning external fixator (Orthofix MultiPlanar MiniRail System). The deformity was corrected gradually until the wrist was in a neutral position, and remained in situ for three months. The external fixator was then replaced with a below-elbow cast to prevent recurrence of the deformity, followed by a wrist gaiter. His hands were splinted and exercised from the time of his admission but we were unable to improve his phalangeal flexion contractures.

The management of his lower limb pathology was focused on his right knee flexion contracture. He was managed in a reverse dynamic sling until his residual knee contracture was fully corrected, after which a cylinder cast was applied, followed by a knee gaiter for long-term use. Mobilisation with a walking frame was gradual due to residual knee pain and weakness.



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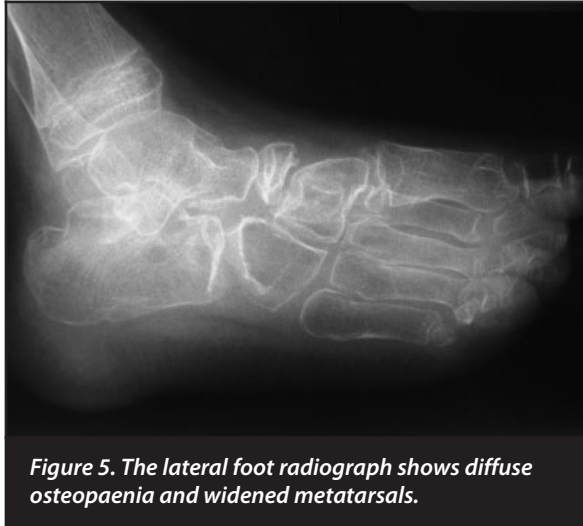


Figure 5. The lateral foot radiograph shows diffuse osteopaenia and widened metatarsals.

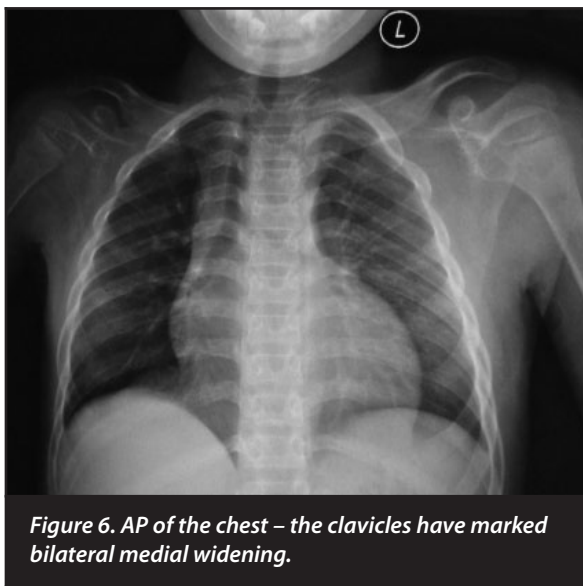


Figure 6. AP of the chest – the clavicles have marked bilateral medial widening.

We assessed his metabolic markers in view of his profound osteopaenia. Dual-energy X-ray absorptiometry showed a Z-score of -3.7SD , and he was subsequently treated with intravenous infusion of a bisphosphonate, zoledronic acid. Radiographs of the distal femur now show 'zebra lines', which have previously been described in association with pamidronate therapy⁸ (Figure 7).

Discussion

The genetic osteolyses are a group of rare disorders which are characterised by radiolucency and progressive disappearance of components of the skeleton. The various forms of osteolysis have been defined and named in terms of the anatomical distribution of the abnormalities, the occurrence of extra-skeletal manifestations and the mode of genetic transmission. There is considerable phenotypic overlap and the syndromic boundaries are blurred. Nosological confusion has been compounded by the inconsistent use of eponymic designations for some forms of osteolysis.

Advances in molecular technology have clarified the situation to some extent and it has become apparent that several apparently disparate entities result from different mutations within the same gene. This observation has led to the concept of a 'spectrum' of overlapping conditions which share a similar biomolecular pathogenesis.

The Torg-Winchester syndrome is a rare autosomal recessive form of osteolysis in which mutations in the matrix metalloproteinase-2 gene (*MMP2*) at the chromosomal locus 16q13 are causative.^{5,6,7,9} The matrix metalloproteinases are a group of enzymes involved in collagen homeostasis. The exact mechanism of how this may lead to the osteolysis syndromes is unknown, but it is believed that abnormal collagen breakdown leads to abnormal osteoblast activity and subsequent osteolysis.³ In this condition the clinical and radiographic features range from osteolysis which predominates in the wrists and ankles (Torg form), to severe generalised osteolysis with skin nodules and a coarse facies (Winchester syndrome).^{2,4} A third condition comprising subcutaneous nodules, arthropathy and osteolysis (NAO syndrome) which has been documented in consanguineous families in Saudi-Arabia is now included in this spectrum.^{1,10} Despite a wide geographical distribution of case reports, the genetic osteolyses are extremely rare. Indeed, in more than 40 years of focus on skeletal disorders in South Africa, only three previous cases have been identified in this country.³

The affected boy whom we have studied has progressive osteolysis with phenotypic manifestations which are all components of the Torg-Winchester spectrum. In particular, predominant but asymmetrical osteolysis of the carpus with widening of the metacarpals, metatarsals and phalanges is suggestive of the former Torg syndrome. The involvement of the elbows and other skeletal regions, together with the joint contractures and muscle wasting are also variable components of this syndrome. Similarly, the severity and distribution of the osteolysis is in keeping with the former Winchester syndrome. Finally, the clavicular widening is characteristic of the NAO syndrome.¹⁰ The fact that the patient has features of all three forms of osteolysis, which were previously regarded as autonomous entities, is entirely in keeping with the concept of a phenotypic spectrum.¹

Differential diagnosis

The most frequent disorders which enter into the differential diagnosis in South Africa include:

- Osteogenesis imperfecta type III
- Bruck syndrome
- Gorham osteolysis
- Polyarticular juvenile idiopathic arthritis
- Juvenile idiopathic osteoporosis

The main condition which enters into the differential diagnosis of the Torg-Winchester NAO spectrum of heritable osteolysis is osteogenesis imperfecta (OI). Diagnostic differentiation of these disorders is especially important in South Africa in view of the relatively high frequency of OI type III in the indigenous African population.¹¹ The manifestations of this autosomal recessive disorder are skeletal fragility, which leads to multiple fractures, together with limb and spinal deformity and marked physical handicap. An unusual complication encountered in a few individuals with this form of OI is the occasional occurrence of flexion contractures of the limbs.

| Table 1: Clinical and radiological features of osteolysis in the affected boy | | |
|--|--|--|
| | Clinical findings | Radiological findings |
| Upper limb | | |
| Right arm and elbow | <ul style="list-style-type: none"> • Atrophy of biceps and triceps muscles • Right elbow fibrous ankylosis deformity, in 90° flexion (<i>Figure 1</i>) | <ul style="list-style-type: none"> • Diffuse osteopaenia • Destruction at elbow joint, which is held in flexion |
| Right forearm and wrist | <ul style="list-style-type: none"> • Atrophy of predominantly the ulnar aspect of forearm | <ul style="list-style-type: none"> • Foreshortened ulna |
| | <ul style="list-style-type: none"> • 45° supination and no pronation present • Ulnar deviation of wrist | <ul style="list-style-type: none"> • Distal ulna articulates with distal radial metaphysis • Carpal bones absent (<i>Figure 2</i>) |
| Bilateral hands | <ul style="list-style-type: none"> • Fixed flexion deformities of little and ring finger interphalangeal joints | <ul style="list-style-type: none"> • Widened metacarpals |
| | | <ul style="list-style-type: none"> • Flexion of ring and little finger interphalangeal joints (<i>Figure 3</i>) |
| Lower limb | | |
| Right knee | <ul style="list-style-type: none"> • Fixed flexion deformity of 80° | <ul style="list-style-type: none"> • Diffuse osteopaenia |
| | <ul style="list-style-type: none"> • No evidence of inflammation | <ul style="list-style-type: none"> • Previous distal femoral growth plate fracture (<i>Figure 4</i>) |
| Bilateral ankles and feet | <ul style="list-style-type: none"> • Limited range of ankle movement | <ul style="list-style-type: none"> • Widened metatarsals and phalanges |
| | <ul style="list-style-type: none"> • Lateral two toes are shortened and flexed at interphalangeal joints | <ul style="list-style-type: none"> • Destruction of great and second toe metatarsophalangeal joints (<i>Figure 5</i>) |
| Chest | | |
| Clavicles | <ul style="list-style-type: none"> • Non-tender | <ul style="list-style-type: none"> • Widening of medial aspects of both clavicles (<i>Figure 6</i>) |

The pathogenesis of this combination, which is termed Bruck syndrome, has not yet been elucidated.^{12,13} The diagnosis of the osteolysis spectrum in the affected boy is strongly supported by the disappearance of the carpal bones, the configuration of the tubular bones of the hands and feet, and the occurrence of significant osteopaenia. An additional radiological discriminant includes the asymmetrical lytic destruction of the affected joints.⁹ In the affected boy, the radiological appearances were entirely in keeping with those of the Torg-Winchester osteolysis as depicted in the definitive *Atlas of Genetic Disorders of Skeletal Development*,¹⁴ and the diagnosis was confirmed by an international expert in the radiology of the osteolyses.¹⁵

Management

There is a paucity of literature concerning the management of the orthopaedic manifestations of the genetic osteolyses. The available literature suggests that the skeletal lysis is progressive, and it is appropriate that the management should focus on correction of deformities and prevention of recurrence.³ Social support by family members is essential to promote rehabilitation.

In the upper limb, the aim of treatment should be to obtain a functional limb to promote independence. Functional assessment by occupational therapists is invaluable in this respect. With regard to the elbow, management will differ depending on the presence of unilateral or bilateral deformities – unilateral deformities should allow functional use of the involved limb, e.g. to bring the

hand to the mouth in order to eat; bilateral deformities should be corrected in order to allow eating and toileting tasks. This can be accomplished by serial casting. With regard to the wrist and hand, the focus is again on functionality – the wrist must attain a neutral position to perform activities of daily living. This can be attained by osteotomy and external fixation with gradual correction, or by serial casting. Phalangeal deformities should be managed with aggressive mobilisation and splinting.



Figure 7. Distal femoral metaphyseal sclerosis subsequent to bisphosphonate therapy

In the lower limb, the focus should be on correction of deformities to allow independent mobilisation. In the event of knee flexion contractures, the correction of the deformity can be obtained using serial casting or reversed dynamic sling traction, an entity that was previously described to correct flexion contractures in haemophiliacs.¹⁶ The ankle if involved must remain plantigrade, as an equinus deformity will prevent mobilisation.

Bisphosphonate therapy

Bisphosphonates have been used in adults with osteoporosis in order to increase bone mass and decrease fracture risk in a broad spectrum of disease. In children, the indications for use in primary and secondary osteoporosis are not as well defined. There appears to be no consensus over the ideal agent, dosage or treatment duration for osteoporosis in children. In this child, initiation of bisphosphonate therapy was based on a low bone mineral density, and was in the form of a zoledronic acid infusion.¹⁷ However, pamidronate has been shown to be ineffective in improving osteolysis or osteoporosis in the appendicular skeleton of individuals with Torg-Winchester syndrome.¹⁸

Radiographs of this child's distal femur exhibit 'zebra lines' which were initially described in association with children on pamidronate therapy. These transverse striations are typical of children receiving cyclical bisphosphonate therapy, and reflect 'increased bone mineralisation, as an indirect response to the inhibition of osteoclastic activity at the time of drug treatment'.⁷ These have been termed 'zebra lines' in order to differentiate them from Harris growth arrest lines, which occur in the metaphyses of rapidly growing long bones following systemic disorders, e.g. malnutrition, femur fractures, infection.⁸ Femoral radiographs also indicate a prior Salter Harris II distal femur fracture, which followed insignificant trauma. This is in keeping with the underlying osteoporotic bone.

Comment

The Torg-Winchester form of osteolysis warrants consideration as a possible diagnosis in children presenting with clinical deformity following minimal trauma and radiological evidence of progressive osteolysis. The orthopaedic management is demanding and includes long-term bracing and night splinting in functional positions to avoid relapses.

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The content of this article is the sole work of the authors. No benefits of any form have been or are to be received from a commercial party related directly or indirectly to the subject of this article. Informed consent was obtained.

Ethical approval for investigation of the heritable connective tissue disorders in South Africa was received by PB from the UCT Health Sciences Faculty Research Ethics Committee ref 026/2010.

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