Acute osteomyelitis in Nigerians with sickle cell disease

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SUMMARY
Acute osteomyelitis comprised 78 (29.3%) of the 266 major skeletal complications seen in 207 patients with sickle cell disease in a five and a half year period. Forty eight (61.5%) of the 78 patients were under the age of 15 years, and the mean age at onset was 12 years (range 9 months to 50 years). Osteomyelitis was often multifocal (in 42% of the cases) and associated with some life threatening disorders. Salmonella accounted for 50% of the 36 organisms isolated from 32 patients with bacteriologically confirmed diagnosis. The 'best guess' antibiotic was a combination of chloramphenicol and cloxacillin. Medical treatment alone proved adequate in most cases. No deaths resulted, but 55% of the patients developed serious complications due partly to the severity of the disease and also to infection involving the epiphyses and joints.

Key words: anaemia (sickle cell), haemoglobin sickle cell disease, Africa (western, Nigeria).

Although acute osteomyelitis is probably the most commonly reported major skeletal complication of sickle cell disease, detailed study of this complication has come mainly from the developed, non-malarial countries (where both sickle cell disease and osteomyelitis are uncommon1); few similar studies have emanated from tropical Africa where sickle cell is very common and skeletal infection rampant.2 3 Previous studies from tropical Africa were either limited to Salmonella osteomyelitis4 5 or they were based on rather few patients.3 6-9

This paper reviews the pattern of acute osteomyelitis in 78 consecutive patients with sickle cell disease seen at the University College Hospital (UCH), Ibadan, in a five and a half year period, and compares it with previous reports from tropical Africa and elsewhere.

Patients and methods
Between June 1976 and December 1981 the author studied prospectively 207 consecutive Nigerians with sickle cell disease and 266 major skeletal complications at the UCH, Ibadan. The major skeletal complications encountered were: osteomyelitis, 129 (48.5%); aseptic necrosis of the femoral head, 75 (28.2%); septic arthritis, 31 (11.7%); pathological fracture, 26 (9.8%); and miscellaneous complications, five (1.9%). Of the 129 patients with pyogenic infection of bone, 75 had acute osteomyelitis; 51 chronic osteomyelitis, and three had both acute and chronic osteomyelitis of different bones. Thus there were 78 cases of acute osteomyelitis, and the present account concerns only these cases.

The initial diagnosis of acute osteomyelitis was clinical, based on history of pains and swelling of the affected part(s), fever, malaise, and inability to use the affected part(s), and on objective signs of pyrexia, localised bone tenderness, and clinical findings suggestive of septicaemia. In most cases the diagnosis was quite obvious and difficulties in distinguishing the disease from bone infarction were infrequent because most of the patients presented late with florid disease.

In 32 patients (41%) the clinical diagnosis of osteomyelitis was supported by a positive blood or pus culture, or both. Twelve patients with negative cultures but with clinical and radiological features consistent with osteomyelitis were presumed to have this disease and were included in the series. No cultures were taken from the remaining 34 cases because of lack of facilities at that time, but these cases were included in the series because they were obviously septic with either discharging sinuses, pus...
and/or persistent toxicity (in the absence of malaria) in the presence of radiological bone changes.

In every case the diagnosis of sickle cell disease was confirmed by haemoglobin electrophoresis and, where possible, by family studies and by estimation of fetal haemoglobin and haemoglobin A2 levels.

**STATISTICAL METHOD**

Student's $t$ test was used to compare means, and the $\chi^2$ test was used to detect association between important characteristics.

**Results**

**AGE, SEX, AND HAEMOGLOBIN TYPE**

The age at onset of osteomyelitis was between 9 months and 50 years (mean 12 years). The mean age for male patients (mean 10 years 3½ months; range 9 months to 23 years) was significantly less than that for female patients (mean 14½ years; range 2 to 50 years) ($p<0.01$). The highest incidence (39 cases or 50%) was in the second decade. Sixty seven patients (85-9%) were under the age of 20. There were 46 male and 32 female patients, a sex ratio of 1.4 to 1. Sixty six patients (85%) had sickle cell anaemia (SS), 11 (14%) sickle cell haemoglobin C (SC), and one had sickle cell thalassaemia.

**KNOWN SITES OF BONE INFECTION**

The total known sites of bone infection were 184. The tibia was the commonest site of involvement. Thirty three patients (29 with SS and four with SC) or 42% had multifocal osteomyelitis. Bilateral symmetrical involvement of the limb bones occurred in nine individuals with SS (Fig. 1), and multiple bones of one side were affected in five cases with SS. Asymmetrical multifocal involvement occurred in nine patients (eight with SS and one with SC). Osteomyelitis of long bones most commonly affected the entire diaphyses, but other sites were not immune. The lower limb was affected almost twice as commonly as the upper limb bones.

**CLINICAL FEATURES**

Most patients had no recognisable predisposing factors; only 16 patients had antecedent infection elsewhere and three others had recent injury. The majority presented late with gross, florid local disease and systemic disturbance (Figs 2 and 3).

Fifty two patients (66-7%) also had associated underlying disorders like septic arthritis (25 patients), severe anaemia with packed cell volume (pcv) less than 20% (18 patients), jaundice (seven patients), pneumonia (seven patients), heart failure (three patients), marasmus (three cases), and septic pericarditis (one case).

**RADIOLOGICAL APPEARANCE**

In 23 patients (29%) initial x rays showed no bone lesion, but subsequent x rays showed obvious bone
changes. Osteolysis and periosteal new bone formation were the commonest bone lesions. Other radiological features included pathological fracture, osteoporosis, and a combination of osteolysis and osteosclerosis.

**BACTERIOLOGICAL FINDINGS**

Thirty-six organisms were cultured from 32 patients (Table 1). Salmonella accounted for 50% of the isolates and gram-negative infections for 80.6%. All but two isolates of *Salmonella* were sensitive to...
chloramphenicol and all but one were sensitive to ampicillin. Twenty six (72%) of the organisms isolated showed in vitro sensitivity to a combination of chloramphenicol and cloxacillin.

OTHER LABORATORY RESULTS
The white cell count (WBC) ranged from 4050 to 124 000/mm³ (4.05–124×10⁹/l) (mean 17 270/mm³ (17.27×10⁹/l)). In 49 patients (62.8%) the WBC was over 11 000/mm³ (11×10⁹/l), while in 34 (43.6%) it was greater than 14 000/mm³ (14×10⁹/l).

The pcv ranged from 5% to 39% (mean 23%). Patients with SS had a significantly lower pcv (mean 22%; range 5–36%) than those with SC (mean 29%; range 13–39%) (p<0.001). Six patients with SS had pcv values between 5% and 14% (mean 10%).

TREATMENT
Initial treatment of critically ill patients comprised correction of dehydration and acidosis, intravenous chloramphenicol and cloxacillin, and packed red cell transfusion and diuretic therapy for those with severe anaemia. Heart failure was treated with digoxin. When the patient was stable the antibiotic, reviewed as necessary, was given orally. Owing to shortage of hospital beds, the less acutely ill patients were treated as outpatients. The duration of antibiotic therapy varied widely but usually lasted for between 12 and 15 weeks. Splintage was an essential adjunct to treatment. Antimalarial therapy and folic acid were used routinely, and analgesics and anti-pyretic drugs were given as necessary.

Thirty eight patients (48.7%) also had some form of operative treatment. Incision and drainage of abscess was performed in 35 bones (30 SS and five SC) of 30 patients (27 with SS and three with SC) and needle aspiration in 11 bones (10 SS and one SC) of eight patients (seven with SS and one with SC).

SEQUELA
There were no deaths, but the incidence and severity of complications were high. In 35 patients (45%) control of infection was achieved without any complication for a minimum of six months. The complications which occurred in the other 43 patients included chronicity (in 20 bones of 17 patients), pathological fracture (in 22 bones of 17 patients), joint stiffness (in eight patients), and septic necrosis of the upper femoral epiphysis in seven patients. Factors that appeared to be associated with chronicity were late presentation at three or more weeks of onset of symptoms; affection of the tibia, femur, or humerus; evacuation of abscess by incision and drainage (in nine patients), and patients aged 10 or more years.

Discussion
The pattern and complications of acute osteomyelitis in sickle cell disease in the present series were similar in many respects to previous reports from West Africa,² ⁴ ⁵ ⁹ but differed in important aspects from reports from non-malarial countries.¹ ¹⁰–¹²

As in previous studies acute osteomyelitis was predominantly a disease of children.¹ ⁴ ¹² The bone most commonly involved was the tibia. In some series, however, the humerus or the femur was the most commonly affected bone.³ ¹³ As in previous reports acute osteomyelitis of long bones was often located in the diaphyses rather than in the metaphyses,¹⁴ even in children, though in an appreciable number the epiphyses were involved, and there was associated septic arthritis in 25 instances. The prognosis is worse with involvement of epiphyses and joints than with diaphyseal osteomyelitis.¹³

Salmonella, the commonest infective organism in the series, accounted for 50% of the positive
cultures. This is consistent with most previous studies from tropical Africa\textsuperscript{3} \textsuperscript{9} and elsewhere.\textsuperscript{1} \textsuperscript{12} \textsuperscript{15} The incidence of \textit{Staph aureus} osteomyelitis in our series was 16-7\%, which is comparable with earlier reports from East Africa.\textsuperscript{3} In contrast, the reported incidence of staphylococcal osteomyelitis in patients without sickle cell disease in Africa\textsuperscript{2} \textsuperscript{3} and elsewhere\textsuperscript{10} is over 50\%. This peculiar association of salmonella osteomyelitis and SS was first reported in 1925,\textsuperscript{16} first recognised in 1951,\textsuperscript{17} \textsuperscript{18} and has been amply confirmed by others.\textsuperscript{2} \textsuperscript{3} \textsuperscript{5} \textsuperscript{15}

From the sensitivity results a combination of chloramphenicol and cloxacillin would be effective against 72\% of the organisms isolated. We therefore propose this combination as the ‘best guess’ antibiotic for acute osteomyelitis in patients with sickle cell disease in West Africa, rather than only chloramphenicol or ampicillin commonly used elsewhere.\textsuperscript{13} We have found this combination to be quite effective in practice. A combination of gentamycin and cloxacillin, which we had earlier suggested should be used in critically ill patients with multifo- cal skeletal infection,\textsuperscript{19} was later found not to be very effective in practice.

The best results as regards control of infection were obtained in the eight patients who were treated by needle aspiration and antibiotics, while the worst results occurred in those treated by open drainage of abscess plus systemic antibiotics. None of the former but nine of the latter patients developed chronicity. Thus needle aspiration should, whenever possible, be the preferred method of evacuating any collection of pus in these patients.

Supported in part by a grant awarded by the University of Ibadan Senate.

References