Plain X-Rays in the Diagnosis of Sickle Cell Limb Pain in Children

A 10-Year Retrospective Audit

ABSTRACT

**Background and aims:** Children with sickle cell disease (SCD) frequently present with limb pain. Differentials include vaso-occlusive episode (VOE) and osteomyelitis (OM). X-rays expose to radiation, but rarely aid in diagnosis. We audited the use of x-ray in investigating children with SCD presenting with limb pain to a South London hospital, and analysed whether x-rays aid in diagnosis.

**Methods:** Patients aged 0-18 years with SCD were identified using the hospital’s SCD database. Admissions from January 2010 to September 2019 in which limb pain was a documented symptom were included.

**Results:** Of 342 patients investigated, there were 188 admissions with limb pain. Diagnoses at discharge were: 174 VOE, 4 OM, and 7 others. 44 (25%) of those with VOE had limb x-rays, compared with 3 (75%) of those with OM. Of those x-rayed, 11 with VOE and all with OM had a subsequent MRI. None of the x-rays assisted in confirming diagnosis or changed management. Of the VOE patients, limb swelling (48% vs 8%, p=<0.0001), fever (57% vs 37%, p=0.021), and peak C-reactive protein (CRP) (109 vs 75 g/L, p=0.044) were determinants of having plan x-rays on admission.

**Conclusions:** X-rays were frequently used to investigate children with SCD but did not appear to influence diagnosis of OM or management. Those with suspected OM should undergo MRIs rather than plain x-rays and be included in new clinical algorithms.

**Keywords:** sickle cell disease, vaso-occlusive crisis, osteomyelitis, X-ray

Frederik Vivian, Subarna Chakravorty*
Department of Paediatric Haematology, King’s College Hospital NHS Trust

*Corresponding author subarna.chakravorty@nhs.net

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Introduction

Sickle cell disease (SCD) is an autosomal recessive blood disorder characterised by frequent episodic pain, increased susceptibility to infection and chronic organ damage. Prevalence in the UK is approximately 1 in 4600, most commonly affecting people of African or African-Caribbean origin. Painful vaso-occlusive episodes (VOE) are the most common cause for admission to hospital. Recurrent VOE can result in bone infarction and osteonecrosis. People with SCD are also at increased risk of osteomyelitis (OM) for reasons including asplenia causing increased susceptibility to encapsulated bacteria, presence of infarcted bone due to osteonecrosis and potential hematogenous spread of gut pathogens due to ischaemia in the gut associated with vaso-occlusion.

It is often difficult to distinguish VOE and OM clinically. Both present with musculoskeletal pain; and swelling, fevers and raised inflammatory markers are common features in both conditions. For a definitive diagnosis of OM, positive blood or operative cultures (such as bone biopsy or joint aspirate) are required. However, blood cultures are often negative despite OM, and bone biopsy is invasive and therefore cannot be used routinely. Radiological evidence can support a diagnosis of OM but plain film x-rays of the limb have a poor sensitivity and specificity for these. Magnetic resonance imaging (MRI) or ultrasound are more likely to guide management, although these can also cause diagnostic uncertainty.

In addition to poor diagnostic yield, plain film x-rays of limbs result in exposure to ionising radiation and is particularly significant in children, who have greater sensitivity to ionising radiation. In patients with SCD, where many presentations with musculoskeletal pain are likely, there is a significant risk from the cumulative radiation dose. Plain x-rays are of little diagnostic benefit in distinguishing VOE from OM and yet they are frequently used as a first line investigation to exclude OM in SCD children presenting with limb pain. An exploration of the effectiveness of plain x-rays in the investigation of limb pain would help future clinical practice.

Method

We undertook an audit of the effectiveness of plain film x-ray to diagnose OM in children with SCD admitted with limb pain to a South London teaching hospital. The retrospective audit covered a period of nearly 10 years between January 2010 and September 2019. Patients with homozygous sickle cell disease and aged 0 to 18 years were identified from the paediatric sickle cell database and were included in the study. The study was approved by our Institutional review board as an audit.

For each admission, data on clinical features, final diagnoses (OM or VOE) and laboratory and radiology results were identified from the hospital electronic records. Results were analysed by final diagnosis and presence/absence of x-ray use.

Statistical significance of differences in clinical features with binomial frequencies (presence of swelling and presence of fever) were calculated using a Chi-squared test. Significance of differences in normally distributed laboratory results was calculated using an unpaired t-test.

Results

A total of 342 patients on our clinical database were analysed with 584 admissions. Of these, 186 admissions had limb pain as a documented symptom, and were included in the audit. Of the 186 admissions for limb pain, the diagnoses at discharge were: 174 vaso-occlusive episodes (VOE), four osteomyelitis (OM), three septic arthritis (SA), two avascular necrosis of the hip, one each of transient synovitis, myositis and non-specific wrist pain.

Imaging

Plain film x-rays were used in 56 of the 186 admissions (30%), with 18 (9.7%) also having MRI scans. Forty-four of the patients admitted for VOE (25%) had x-rays. Seven patients with VOE, all patients with OM and SA, went on to having MRIs. None of the x-rays appeared to have contributed to the final diagnosis or changed management.

Vaso-occlusive episodes
Patients presenting with limb pain and a diagnosis of VOE were more likely to be investigated with limb x-rays if they had swelling of the affected limb (48% vs 8%, p=<0.001), fever (57% vs 37%, p=0.021) and an elevated mean C-reactive protein level (109 vs 75 g/L, p=0.044), with no significant difference in total white cell count.

**Osteomyelitis**
All patients with OM presented with localised swelling and three out of four presented with fever. Peripheral blood cultures were sent in 45% of patients with VOE, all patients with OM and SA. Of the 45% of VOE patients where blood culture were taken, 3 had positive results growing organisms that are frequent contaminants [Table 2]. Only one of the patients with OM had a positive peripheral blood culture result while another was positive from a bone biopsy.

**Pain**
All patients with OM had unilateral limb pain, compared with VOE patients where the median was pain affecting more than 2 sites. The number of painful sites was not found to be a determinant of who had x-rays. Patients with OM had a longer duration of pain prior to admission, compared to those with VOE (8 vs 2.47 days) [Table 3].

**Antibiotics**
In this audit, 97 children (56%) with a diagnosis of VOE received broad spectrum antibiotics for a mean duration of 7 days, with some of these having alternative indications for antibiotic treatment.

**Discussion**
While OM is a serious diagnosis in children with SCD, it is rare compared to vaso-occlusive pain mimicking OM. X-rays do not appear to aid in this diagnostic challenge, due to low sensitivity for both conditions and lower specificity. MRI scans can be helpful, and are an important investigation in OM, although infarcted and infected bone can appear similar in this modality. Positive blood cultures with pathogenic bacteria improves confidence of a diagnosis of OM, but the sensitivity of blood cultures is relatively low. Culture of an operative biopsy sample is a definitive investigation to confirm OM, but due to its invasive nature, is not appropriate unless there is a very high clinical suspicion. The investigation of children with bone pain and infective symptoms is therefore a difficult area for paediatricians and haematologists.

In our analysis of admissions for limb pain, 25% of patients with VOE had a plain x-ray. Plain x-rays offered no benefit to the diagnostic process. Those patients presenting with features suggestive of OM such as an elevated CRP, fever or limb swelling were likely to have a plain x-ray as their initial investigation. In none of the 186 admissions, did a limb x-ray aid diagnosis or alter management. Thus in such cases of suspected OM, an MRI of the affected limb would be a more appropriate investigation.

Children with suspected OM are treated with broad spectrum empirical antibiotics, and many of these are discontinued once clinical suspicion of OM reduces during the admission episode. There appears to be variation in institutional practices in the use of empirical antibiotics in SCD children presenting with limb pain. In this audit, 56% of children received broad spectrum antibiotics compared to 86% of 358 VOE patients in another paediatric cohort from London.

In this audit we examined admissions of patients on our current SCD paediatric database over a 10-year period. This therefore excluded admissions of patients that occurred within the last 10 years, but that are now older than 18 and are therefore no longer in our current database. This resulted in an age-skew of admissions, with more admissions of younger patients. While VOE commonly presents from a young age onwards, OM is more likely in older children. Since the purpose of the audit was to examine the plain x-ray use in children presenting with limb pain, we chose to study all children with pain, rather than identify those with proven or probable OM and work retrospectively to their original presentation. A longer study looking at the clinical features and investigation variation between VOE and OM may offer greater insight into how these can be distinguished and is an area that would benefit from further research.
We have found a high frequency of x-ray (30%) use for investigation of limb pain in children with sickle cell disease resulting perhaps in unnecessary irradiation in a population at risk of significant lifetime exposure. Evidence based clinical algorithms or guidelines for suspected OM incorporating features such as fever, unilateral / single site of pain and raised inflammatory markers may help target appropriate imaging such as MRI and reduce the use of plain x-rays in the management of limb pain in children with SCD is needed. We plan to re-audit the use of plain x-rays once these measures are put in place and hope to achieve a reduction in x-ray use in this group of children.

References

Table 1: Frequency and percentage of plain film x-ray use, by final diagnosis

<table>
<thead>
<tr>
<th>Final diagnosis at discharge</th>
<th>Number</th>
<th>Frequency of x-ray use (percentage)</th>
</tr>
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<tbody>
<tr>
<td>VOE</td>
<td>174</td>
<td>44 (25)</td>
</tr>
<tr>
<td>OM</td>
<td>4</td>
<td>3 (75)</td>
</tr>
<tr>
<td>Septic arthritis</td>
<td>3</td>
<td>2 (67)</td>
</tr>
<tr>
<td>Myositis</td>
<td>1</td>
<td>1 (100)</td>
</tr>
<tr>
<td>AVN</td>
<td>2</td>
<td>2 (100)</td>
</tr>
<tr>
<td>Transient synovitis</td>
<td>1</td>
<td>1 (100)</td>
</tr>
<tr>
<td>Wrist pain</td>
<td>1</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Total</td>
<td>186</td>
<td>56 (30)</td>
</tr>
</tbody>
</table>
### Table 2: Organisms isolated from blood culture, by final diagnosis

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Organisms isolated (from blood cultures unless stated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VOE</td>
<td><em>Staphylococcus caprae, Staphylococcus epidermidis, Streptococcus viridans</em></td>
</tr>
<tr>
<td>OM</td>
<td><em>Abiotrophia defectiva, Salmonella (negative blood cultures, positive intra-operative sample)</em></td>
</tr>
</tbody>
</table>

### Table 3: Clinical features and investigations, by final diagnosis

<table>
<thead>
<tr>
<th></th>
<th>VOE (all)</th>
<th>VOE (≥1 limb x-ray)</th>
<th>VOE (no limb x-ray)</th>
<th>OM</th>
<th>SA</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>174</td>
<td>44</td>
<td>130</td>
<td>4</td>
<td>3</td>
<td>5</td>
</tr>
</tbody>
</table>
| Number (%) females        | 57 (33%)  | 13 (30%)            | 44 (34%)            | 3  | 1  | 2 (40)
| Number (%) on hydroxycarbamide | 93 (53%)  | 27 (61%)            | 66 (51%)            | 2  | 1  | 2 (40)
| Mean age at admission, years (s.d.) | 8.26 (4.7) | 7.45 (5.6) | 8.54 (4.4) | 14.5 (1.7) | 7.33 (3.5) | 5.6 (5.5)
| Limb x-ray used            | 25%       | 100%                | 0%                  | 75%| 66.7%| 80%
| MRI used                   | 7%        | 25%                 | 0.8%                | 75%| 66.7%| 40%
| Peak WCC, mean (s.d.)      | 14.8 (8.0)| 16.6 (12.3)         | 14.2 (5.8)          | 12.3 (3.0)| 22.3 (2.1)| 14.4 (6.7)
| Peak CRP, mean (s.d.)      | 84.3 (95.3)| 113.2 (90)         | p=0.044*            | 74.5 (95.5)| 77.5 (111.1)| 130.3 (16.9)| 68.2 (106.6)
| Swelling present           | 18%       | 48% (p<0.001)*     | 8%                  | 100%| 100%| 40%
| Fever present              | 42%       | 57% (p=0.021)*     | 37%                 | 75%| 67% | 0%
| Max. height of fever, ºC, mean | 38.4 | 38.3                 | 38.5                | 38.5| 38.4| n/a
| Blood Cultures sent         | 45%       | 62%                 | 39%                 | 100%| 100%| 20%
| Positive Blood Cultures     | 3         | 1                   | 2                   | 1  | 0  | 0     |
| Biopsy / Aspiration         | 0.57%     | 2.27%               | 0%                  | 50%| 50%| 0%
| Mean number of sites affected (s.d.) | 2.5 (1.2) | 2.3 (1.3)            | 2.6 (1.2)           | 1 (0)| 2.0 (1) | 1.2 (0.4)
| Mean duration of pain prior to admission, days (s.d.) | 2.5 (3.2) | 3.2 (4.6)            | 2.2 (2.5)           | 8 (9.0)| 3.3 (3.2) | 19.2 (39.6)

OM= osteomyelitis, SA= septic arthritis, s.d.= standard deviation, WCC= white cell count, CRP= C-reactive protein

*Patients with VOE who received x-rays were more likely to have higher mean CRP (113.2 vs 74.5, p=0.44), swelling of the affected limb (48% vs 8%, p<0.001) and fevers (57% vs 37%, p=0.021)