Melorheostosis of The Leg: A Case Report

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ABSTRACT
Melorheostosis is a benign hyperostotic disease of the peripheral skeleton, rarely involving the axial skeleton. This disease is associated with ossified and non-ossified soft tissue masses surrounding the joints. We report the case of a 28-year-old male who presented to an orthopedic clinic with a chronic history of right leg pain. Radiological evaluation using X-ray, computed tomography, and magnetic resonance imaging showed features consistent with that of melorheostosis. Recognition of this entity by clinicians can avoid unnecessary investigations and biopsy.

CASE REPORT
A 28-year-old male with no history of medical illness or previous trauma presented to the orthopedic clinic with mild right leg pain that persisted for 6 months. The patient described his pain as a dull-ache accompanied with a tingling sensation to touch.

The results of the clinical examination were unremarkable apart from mild tenderness and numbness extending along the anterior lateral aspect of the right leg. Vital signs were all within normal ranges, and no muscle weakness or limitation of joint mobility was observed. Lab tests, including complete blood count (CBC), erythrocyte sedimentation rate (ESR), and C-reactive protein (CRP), were normal.

For further evaluation, the patient was referred to the radiology department. Anteroposterior and lateral radiographs of the right leg were initially obtained. Results revealed sclerotic changes in the proximal two-thirds of the fibula with periosteal and endosteal cortical thickening, consistent with the characteristic “dripping wax” appearance of melorheostosis (Figure 1). Magnetic resonance imaging (MRI) and computed tomography (CT) were subsequently performed to define the extent of pathology and to exclude associated complications. MRI results demonstrated a low signal intensity lobulated perosteal new bone formation involving the posterolateral cortex of the right fibular shaft. Clear endosteal thickening with no medullary invasion or soft tissue masses was observed (Figure 2). CT scan results showed a lobulated undulating perosteal and endosteal thickening partially encroaching the medulla, corresponding to the low signal abnormality observed on the MRI. No fracture was identified (Figure 3).

The patient was treated symptomatically; he was prescribed nonsteroidal anti-inflammatory drugs for 1 month and has been followed at the orthopedic clinic with partial improvement of symptoms.

DISCUSSION
Etiology & Demographics:
Melorheostosis, also known as Leri disease and candle bone disease, is a benign hyperostotic disease [1]. It is a rare condition with an estimated prevalence of 0.9 per million [2] and affects both sexes equally [3]. The disease manifests during late childhood or early adulthood; however, it can occur...
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Musruloskeletal Radiology:

at any age [1,3]. It is a unilateral disease of the peripheral skeleton, more commonly affecting the lower extremities with a diaphyseal predilection [2]. Melorheostosis rarely affects the axial skeleton [1].

The exact etiology of melorheostosis remains unclear [1]; however, there are two major theories attempting to explain its origin. Murray and McCredie hypothesized that melorheostosis results from insult to segments of the neural crest during embryogenesis, resulting in segmental sensory lesions. This hypothesis is based on the observation that melorheostosis correlates with sclerotomes in their distribution [4]. Conversely, Fryns hypothesized that melorheostosis results from postzygotic mutations in the mesenchyme. These mutations lead to variable and asymmetric involvement of the skeletal structures, and vascular and hamartomatous changes in the overlying soft tissues [5].

Despite the increased prevalence of melorheostosis in families with autosomal dominant osteopoikilosis the disease is nonhereditary [1].

Clinical & Imaging Findings:

Patients with melorheostosis are usually asymptomatic; however, various symptoms, including pain, limitation of movement, deformities, limb swelling, and limb length discrepancy may occur in growing children [1]. In most patients seeking medical attention, pain is usually the presenting symptom [6]. Melorheostosis may be associated with ossified and non-ossified soft tissue masses, usually surrounding the joints. The symptoms may arise due to impingement on nearby structures, either directly or through associated soft tissue masses. These masses are not necessarily a continuum of the disease lesions [1]. Clinical course may fluctuate with alternating exacerbation and remission periods [1]. Melorheostosis has been associated with various conditions, including neurofibromatosis, tuberous sclerosis, linear scleroderma, tricho-dento-osseous syndrome, rheumatoid arthritis, hypophosphatemic rickets, vascular malformations, and hemangioma [3].

The disease may extend along the affected cortex in a proximal to distal pattern, resembling the pathognomonic “dripping candle wax” appearance on an X-ray and CT scan. Other radiologic appearances include osteoma, an osteopatia striata-like appearance, and myositis ossificans-like lesions without lamellae in soft tissues [1]. CT scans show cortical thickening that might extend to the medulla, with a clear demarcation between the normal bone and the hyperostotic lesions [3,7]. Additionally, CT scans clearly depict soft tissue abnormalities associated with the disease. MRI typically shows cortical hyperostosis of low signal intensity on all pulse sequences [1]. Rarely, MRI may show enhancing soft tissue masses that resemble aggressive neoplastic lesions [8]. Melorheostosis typically show high uptake of the tracer on bone scintigraphy that crosses the adjacent joints [3].

Treatment & Prognosis:

Many surgical and nonsurgical treatment options exist; however, all treatment options are symptomatic rather than curative, and are directed toward the disease symptoms and complications [1]. Therapy is primarily aimed at relieving the pain and restoring full range of motion. Examples of treatment modalities include analgesia, manipulation, braces, serial casting, physiotherapy, nerve blocks, and sympathectomy [3]. Patients experiencing mechanical symptoms, such as nerve compression, contractures, impingement, and deformity should be offered surgical alternatives [6]. Symptomatic cases are eligible for surgical excision [9]; however, the recurrence of disease remains a possibility [6]. Although the condition is not fatal, it causes considerable morbidity. Melorheostosis typically has a benign course, however two case reports of malignant transformation to osteosarcoma and malignant fibrous histiocytoma have been reported in literature[15].

Differential Diagnoses:

Melorheostosis has to be differentiated from common differential diagnoses, including but not exclusive to myositis ossificans, osteoid osteoma, parosteal osteosarcoma, osteochondroma, and chronic osteomyelitis [7].

Myositis Ossificans:

Myositis ossificans is a condition stemming from trauma or an injury. Radiographs show peripheral ossification with central luency. The lesion is usually distant from adjacent bone. CT scans demonstrate peripheral calcification early in the disease course before it becomes evident on conventional radiographs.

MRI scans show iso- or slight hyperintensity within the lesions on T1-weighted and T2-weighted images. Diffuse or rim enhancement may also be observed [10].

Osteoid Osteoma:

On radiographs, intracortical radiolucency, known as intracortical nidus, surrounded by cortical thickening can be observed. The nidus is more apparent on CT scans and may show internal calcification. On MRI, the nidus appears to have low to intermediate signal intensity on T1-weighted images and variable signal intensity on T2-weighted images. Inflammatory changes surrounding the bone marrow and soft tissues are better observed on MRI [11].

Parosteal Osteosarcoma:

Plain radiographs show a mushroom-like mass with a broad base attachment protruding from the underlying cortex. The connection between the tumor and underlying bone cortex can be well visualized using CT and MRI. Periosteal reaction is commonly minimal, and the underlying bone cortex may be thickened or partially eroded. CT is the most accurate modality for evaluating the underlying cortical bone integrity, and MRI is better for demonstrating bone marrow invasion [12].
**Osteochondroma:**
On radiographs, the cortex of the lesion appears to be attached to the cortex of underlying bone, and the medullary cavity of the lesion is continuous with the medullary cavity of the underlying bone. In addition, the lesion is capped with cartilage. In comparison with radiographs, CT scans delineate the connection between the cortex and the medullary cavity of the lesion with the original bone in a better manner than radiographs. The best modality for measuring cartilage cap thickness is MRI because it shows high signal intensity on T2-weighted images [13].

**Chronic osteomyelitis:**
In chronic osteomyelitis, a sequestrum is frequently visible on radiographs as focal sclerosis surrounded by a lucent rim. In some cases, the sequestrum becomes encased in a thick sheath of new bone known as an involucrum. The bone cortex may appear to be destructed. These findings are better visualized on a CT scan. MRI shows a dark sequestrum in all sequences surrounded by hypervascular granulation tissue with post-contrast peripheral enhancement. The involucrum either displays a normal signal or edema. A defect, known as cloaca, connecting the medulla to the surrounding soft tissues may be visualized. Draining pus within the cloaca displays high signal [14].

**TEACHING POINT**
Melorheostosis is a rare, often asymptomatic disease; however, it warrants consideration and a clear differentiation from more deleterious abnormalities in patients presenting with chronic limb pain. Melorheostosis is characterized by classic signs on radiographs and CT such as the “dripping candle wax”. The clinician can avoid unnecessary investigations and biopsy if asymptomatic, while management depends on various particular clinical presentations or complications.

**REFERENCES**


Figure 1: 28-year-old male with melorheostosis.
Findings: (a, b) Anteroposterior and lateral radiographs of the right leg show sclerosis on the posterior lateral aspect of the proximal two-thirds of the fibula with periosteal and endosteal cortical thickening, consistent with the classic dripping wax appearance (arrows).
Technique: X-ray of tibia/fibula, kVp 45, mAs 4.

Figure 2: 28-year-old male with melorheostosis.
Findings: Sagittal T1-weighted (Figure 2a), axial T1-weighted (Figure 2b), axial T2-weighted with fat suppression (Figure 2c). MRI demonstrates low signal intensity lobulated periosteal new bone formation involving the posterolateral cortex of the right fibular shaft (arrows). Clear endosteal thickening is best observed on axial images (Figure 2b-c). No medullary invasion or soft tissue masses were evident.
Technique: MRI right leg without contrast (GE 1.5T Signa)
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Musculoskeletal Radiology:

Item | Melorheostosis
--- | ---
Etiology | Unknown
Prevalence | 0.9 per million
Sex ratio | Male to Female 1:1
Age predilection | It can occur at any age; however, is most prevalent in late childhood to early adulthood
Risk factors | No identified risk factors
Treatment | Symptomatic
Prognosis | Non-fatal disease with considerable morbidity
Findings on imaging | X-ray: candlewax appearance
CT scan: candlewax appearance with high density cortical thickening
MRI: cortical hyperostosis of low signal intensity on all pulse sequences.

Table 1: Summary table of melorheostosis.

Figure 3: 28-year-old male with melorheostosis.
Findings: Selected axial (Figure 3a) with reconstructed coronal (Figure 3b) and sagittal (Figure 3c) images of a non-contrast computed tomography of the leg show lobulated undulating periosteal and endosteal thickening partially encroaching the medulla, corresponding to MRI low signal abnormality (arrows). No fractures are evident.
Technique: MDCT (GE Discovery CT750 HD), 120 KVP, 149 mA, 0.6-mm slice thickness.
<table>
<thead>
<tr>
<th>Melorheostosis</th>
<th>Classic dripping candlewax appearance</th>
<th>Candlewax appearance with high attenuation cortical thickening</th>
<th>Cortical hyperostosis of low signal intensity on all pulse sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myositis ossificans</td>
<td>Peripheral ossification with central lucency and clear demarcating line between the lesion and normal bone</td>
<td>Peripheral calcification of the lesion becomes evident earlier on CT than on conventional radiographs</td>
<td>Iso- or slight hyperintensity on T1W and T2W images, respectively; diffuse or rim enhancement may also be observed</td>
</tr>
<tr>
<td>Osteoid osteoma</td>
<td>Intracortical nidus surrounded by cortical thickening</td>
<td>The nidus is more apparent on CT than radiographs with low or high attenuation</td>
<td>The nidus appears to have low to intermediate signal intensity on T1W images and variable signal intensity on T2W images</td>
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<td>Parosteal osteosarcoma</td>
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<td>Osteochondroma</td>
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<td>Usually shows focal sclerosis surrounded by a lucent rim, known as a sequestrum; it also shows the involucrum that surrounds the sequestrum</td>
<td>The findings in conventional radiographs are better visualized by CT scan</td>
<td>MRI shows a dark sequestrum in all sequences surrounded by hypervascular granulation tissue with post-contrast peripheral enhancement; cloaca may be visualized; draining pus within the cloaca displays a high signal</td>
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</tbody>
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Table 2: Differential diagnoses table for melorheostosis.

**ABBREVIATIONS**

AP = Anterior posterior  
CBC = complete blood count  
CRP = C-reactive protein  
CT = computed tomography  
ESR = erythrocyte sedimentation rate  
KVP = Peak Kilovolt  
mAs = milliamperes seconds  
MDCT = Multidetector computed tomography  
nm = millimeter  
MRI = magnetic resonance imaging  
T1W = T1-weighted  
T2W = T2-weighted

**KEYWORDS**

Melorheostosis; Leri disease; Candle bone disease; Fibula; Magnetic resonance imaging; MRI

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