Spleen uptake on bone scan

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Abstract

We present a patient with spleen uptake on bone scan that was due to sickle cell disease. We also discuss other etiologies for this finding.

Case Report

A 24-year-old female with a past medical history of sickle cell disease presented to the hospital complaining of right hip pain after a fall at home. Diagnostic imaging including CT head, hip X rays, thoracic and lumbar spine X rays were unremarkable. As she continued to complain of right hip pain, a bone scan was ordered.

Following the intravenous administration of 877 Mbq Tc-99m MDP, flow and blood pool images of the pelvis were obtained (not shown), followed by delayed whole body and spot views of the pelvis. Whole body images demonstrate diffuse increased radiotracer uptake in the left upper quadrant of the abdomen just superolateral to the left kidney, suggesting uptake in the spleen (Fig 1). Diffuse increased uptake throughout the skull, distal femurs and proximal tibias is in a pattern characteristic of bone marrow hyperplasia.
FIGURE 1.

Anterior (left) and posterior (right) bone scans show unexpected uptake in the left upper quadrant characteristic of the spleen (black and white arrow). There is also increased radiotracer uptake in the skull, distal femurs and proximal tibias (black arrows) characteristic of the pattern of bone marrow expansion.

The peripheral blood smear showed normocytic-normochromic anemia with sickle cells, Howell-Jolly bodies and Pappenheimer bodies. The patient also had a history of iron overload from multiple transfusions and was on chelation therapy.

Pappenheimer bodies (Fig 2) are blue-purple inclusion bodies located within the red blood cells formed by phagosomes that have engulfed excessive amounts of iron. They are present in diseases such as sideroblastic anemia, hemolytic anemia, and sickle cell disease.

Howell–Jolly bodies (Fig 2) are basophilic nuclear remnants (clusters of DNA) in circulating erythrocytes. These inclusions are normally removed by the spleen but persist in individuals with functional hyposplenia or asplenia. Common causes include asplenia (post-splenectomy), trauma to the spleen, and autosplenectomy caused by sickle cell anemia.

Discussion
The most common causes of uptake in the spleen on the bone scan are hemosiderosis, hemochromatosis and sickle cell disease. Some of the uncommon causes of splenic uptake are neoplastic deposits to the spleen (e.g., from breast ca, Hodgkin’s disease), splenic hemangioma, splenic artery calcification, frequent platelet and RBC transfusions (1) and thalassemia major. Rare causes of splenic uptake include Glucose -6-phosphate dehydrogenase deficiency (G6PD deficiency).

Hepatic and splenic uptake of Tc-99m MDP has also been reported when MDP is injected within a short interval or up to 2 days after the administration of gadolinium-DTPA MRI contrast agents and even when MDP is injected before Gd-DTPA (2,3). Another cause of splenic uptake could be a very recent prior radionuclide study, such as with $^{99m}$Tc-sulfur colloid or $^{111}$In-labeled white blood cells. Both will additionally demonstrate radiotracer uptake in the liver.

The mechanism of splenic uptake of bone-seeking agents in sickle cell anemia has been presumed to be deposition of calcium. Microscopic deposits of calcium that could not be detected radiographically might also cause uptake of Tc-99m MDP. Several ultrastructural studies have demonstrated mitochondrial binding of calcium ions in necrotic muscle cells, within a crystalline structure similar to that of hydroxyapatite (4).

Hemosiderosis occurs as a result of recurrent transfusion and increased deposition of iron from the sequestration of abnormal RBCs within the spleen. Several studies including a study by Jones et al (5) proposed that iron deposits might be responsible for uptake of bone-seeking radionuclides. The shortened survival of the abnormal sickle cells leads to a compensatory marrow expansion throughout the skeleton. An expanded marrow space results in increased uptake of tracer in the calvarium, periarticular locations in the long bones, and even the distal extremities.

**Conclusion**

It is not uncommon to see incidental uptake of bone seeking agents in the extraosseous structures, e.g., spleen, liver, breast and lungs. Review of patient’s history, laboratory findings and other imaging work up provides useful clues in reaching the diagnosis.
Spleen uptake in the bone scan can be caused by number of diseases including Sickle cell disease as discussed in our case report. It is presumed to be caused by microscopic or macroscopic splenic calcifications or prior infarcts. Iron overload in patients receiving blood transfusions can be another cause.

References


