THE CASE

A 22-year-old man presented to our family medicine clinic with hip pain of 2 weeks’ duration. The patient played hockey around the time of onset, but denied any specific injury. The pain, which affected the anterolateral aspect of the patient’s right hip, first started when he stood up after eating a meal. He rated the pain as an 8/10 on average and said that it was worse with movement. The patient had not shown improvement with conservative therapy (rest, ice, and ibuprofen). His medical and surgical history were noted as noncontributory. He was not taking any medications other than over-the-counter pain medication, did not drink alcohol or use tobacco, and he exercised regularly. A review of systems was negative except for right hip pain.

The physical exam revealed pain on active flexion and abduction of the hip. Passive range of motion (ROM) was negative for pain. The right hip was grossly normal with no pain on palpation or crepitus. There was no associated muscle tenderness. The patient was advised to continue to rest and ice the hip, as well as to take ibuprofen for pain relief. He was referred to Physical Therapy.

He returned to our clinic 4 weeks later with no improvement in his symptoms despite several sessions of physical therapy. We ordered radiographic images and magnetic resonance imaging (MRI) of the right hip.

THE DIAGNOSIS

Plain films (FIGURE 1A) showed bilateral avascular necrosis (AVN) of the femoral heads, which was worse on the right side than the left. An MRI (FIGURE 1B) further supported this diagnosis, revealing changes in the femoral neck consistent with a stress reaction and no significant collapse of the femoral head.

DISCUSSION

AVN of the hip has an incidence ranging from 10,000 to 20,000 new cases annually.1,2 It has many possible causes, including trauma, systemic lupus erythematosus, glucocorticoid use, and chronic excessive alcohol use. Although the underlying pathophysiology varies, experts hypothesize that most cases are caused by a disruption of the blood supply, which leads to hyperemia and cortical destruction and collapse.1,2

Certain medications can cause AVN

A more thorough history-taking at this patient’s initial visit would have prompted imaging at that time and ensured that the standard of care was met. Upon further investigation at his follow-up appointment, it was discovered that he had been diagnosed with acute pre-B cell lymphoblastic leukemia (ALL) 2 years earlier and had undergone chemotherapy with cytarabine, vincristine, L-asparaginase, daunorubicin, methotrexate, and glucocorticoids. This discovery,
along with the lack of symptom improvement, prompted the ordering of his imaging studies. Long-term glucocorticoid therapy is the second leading cause of AVN, following traumatic events. High daily dosages (>40 mg/d) and high cumulative dosages of glucocorticoids are associated with a significantly increased risk for AVN.4,5

The other chemotherapy agents with which our patient had been treated (cytarabine, vincristine, L-asparaginase, daunorubicin, and methotrexate) have no reported links to AVN. When mentioned in the literature, however, they are usually coupled with the use of dexamethasone or prednisone.

One case report described a patient with acute promyelocytic leukemia who was treated with all-transretinoic acid, daunomycin, cytarabine, and a short course of dexamethasone, and was diagnosed with AVN 2 years after the cessation of chemotherapy.6 This demonstrates that steroid use does not need to be recent to have a contributory effect.

Did leukemic burden play a role?
We also considered whether the patient’s leukemic burden contributed to his osteonecrosis. Leukemia and its therapy regimens have been reported to cause cerebrovascular complications,7 so it would be logical to postulate that they might also pose a risk to the vasculature of the femoral head. One case report describes hip pain and AVN as the initial manifestation of chronic myeloid leukemia (CML).8 But CML is more often associated with a severely increased white blood cell (WBC) count than is ALL, and our patient’s WBC count was in the expected range for a patient in the maintenance phase of chemotherapy, making leukemic burden a less likely culprit.

Know your patient’s history
Our patient had received an initial dose of approximately 120 mg/d prednisone alone during the first 28 days of his induction therapy for ALL. In addition, he received dexamethasone maintenance therapy, which can accumulate to >140 mg/m² over the course of therapy.9 This information was ultimately integral to his diagnosis and treatment.

Our patient was referred to Orthopedics. He underwent therapy with alendronate and did not require surgical intervention.

THE TAKEAWAY
This case illustrates the importance of obtaining a thorough medical history, including
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previous drug exposures, as a means to raise or lower one's index of suspicion appropriately.

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References