Sir,

Pediatric and adolescent ALL cases presenting with extensive lytic bony lesions with or without hypercalcemia are anecdotal and hence may prove baffling to the investigating physicians and diagnosis may be delayed, especially when there are no blasts in the peripheral blood film. However, presence of unexplained anemia and hypercalcemic symptoms can be warning clues to an astute clinician. Description and short course of one such case is presented.

A 5 years male child presented to the department of Pediatrics with complaints of anorexia, abdominal pain, constipation, polyuria, polydipsia, bony aches and pains of 2 months duration. In addition, there was history of fever, irritability, nausea and vomiting. The child was initially managed at maternity and child health service and later, shifted to our ward for evaluation of anemia. On examination, he was sick looking, conscious and well-oriented, febrile and pale. Bilateral cervical lymph nodes were enlarged multiple, about 1x1 cm. Bones were tender and bilateral hip joint movements were painful and restricted. Rest of systemic examination was within normal limits. Investigations Hb 7.7 g/dl, WBC; 6700/cmm. ESR 30mm/hr, differential No blasts. Platelets-268/cmm. Liver and renal function tests were normal. Serum uric acid 10.6 mg%. LDH 1330 i. u/l. Serial serum calcium and phosphorus readings were - 17.84, 13.0, 11.26 mgms/dl and 4.9, 4.9, 4.02mg/dl, respectively. 24 hour urinary calcium was 630 mgms (normal upto 300 mg/day). Electrocardiogram, cerebrospinal fluid and urine analysis were within normal limits. chest x-ray showed normal lung fields, x-ray skull and pelvis revealed diffuse lytic lesions. Abdominal USG scan normal. Plasma PTH - 3 pg/ml (n, 10-69 pg/ml). In view of low Hb, raised Ca ++, lytic lesion and low PTH, a hematological disorder was considered. Bone marrow aspiration showed 44% blasts. Blasts were PAS +ve, SB/MPO – negative and were positive for CD3, CD20 and CD45- 44.46%. A diagnosis of ALL-L2 was made. Patient was started on UKALL-12 protocol and on subsequent follow up, patient was asymptomatic, bone pain had relieved, lytic lesions regressed, and remission was induced. Hypercalcemia, uric acid and serum lactate dehydrogenase levels normalized.

COMMENTS:

Bone involvement is common in ALL. About one third patients present with bone pains and approximately one-half of these have bone involvement. Presence of frank osteolytic lesions with hypercalcemia is infrequent. Bone destruction may be the cause of hypercalcemia besides other factors. Bony lytic lesions with accompanying hypercalcemia can develop at the onset. The hypercalcemia in pediatric ALL patients with early pre-B cell was estimated to be about 4.8% in a series of 83 cases from Japan with PTHrP levels lever elevated to 112-240 pmol/l (normal range is 17.61.2 pmol/l) in all. Normalization of both Increased plasma prostaglandin E 2 (PGE2) levels and hypercalcemia can occur after chemotherapy. Presentations as in our case have also been reported in adult T cell leukemia as well as in adult T cell leukemia/lymphoma. In the latter, it may be associated with elevated levels of interleukin-6 and PTHrP. Hypercalcemia in cancer can be due to release of tumour necrosis factor (alpha and beta), IL1, 2, 6, TGF beta, 1, 25(OH) 2 and direct invasion,
apart from excess of PTHrP\textsuperscript{10} and PGE2 and rarely due to PTH secretion. The later may be secreted directly by the lymphoblasts.\textsuperscript{11} Frequently, blasts are absent from the peripheral blood in such cases.\textsuperscript{4,7,12} A single patient of acute megakaryocytic leukemia (M7) presenting with hypercalcemia and skeletal lytic lesions has also been described.\textsuperscript{13} The final outcome of children with hypercalcemia is similar to children with ALL indicating that the development of hypercalcemia itself is not a poor prognostic factor.\textsuperscript{14} Curiously such manifestations, have rarely been reported in CML, CLL in adults. Apart from treatment of the basic disease, hypercalcemia needs to be addressed with measure like saline diuresis, calcitonin and bisphosphonates.

Intriguingly, apart from anemia peripheral blood counts were maintained in this patient also and he presented chiefly with clinical features of hypercalcemia at the onset and had a delay in diagnosis. Hence, in a child presenting with a short illness with hypercalcemia and anemia, ALL should be included the list of differential diagnosis. Furthermore, rising serum calcium levels after normalization could herald a leukemia relapse in such a case...

REFERENCES:


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