Technical Note

Long segment spinal epidural extramedullary hematopoiesis

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Abstract

**Background:** Extramedullary hematopoiesis is defined as the formation of blood cells outside the bone marrow. It is a common manifestation of many chronic hemolytic anemias, and typically involves the liver, spleen, and lymph nodes. Only rarely is the spinal epidural space involved.

**Methods:** We describe a 25-year-old male, known to have thalassemia intermedia, who presented with a 1-month history of stiffness and weakness in both lower extremities. On physical examination, he had palpable splenomegaly accompanied by spinal tenderness at the D5 level, weakness in both lower extremities, hyperactive bilateral Patellar and Achilles reflexes with bilateral Babinski responses, and a graded sensory loss to pin appreciation below D5.

**Results:** The magnetic resonance (MR) study revealed a posterior, isointense and soft tissue epidural mass extending from D2 to D12 on both the T1- and T2-weighted images. These findings were consistent with the diagnosis of “red marrow,” and long-segment spinal epidural extramedullary hematopoiesis.

**Conclusions:** Although extramedullary hematopoiesis is rarely encountered within the spinal canal, it should be considered among the differential diagnoses when a posterior compressive thoracic lesion contributes to myelopathy in a patient with a history of thalassemia intermedia and the accompanying chronic hemolytic anemia.

**Key Words:** Extramedullary hematopoiesis, epidural, thalassemia

INTRODUCTION

Although thalassemia, a form of chronic hemolytic anemia, is commonly associated with extramedullary hematopoiesis (EMH), it is only rarely encountered in the spinal epidural space. Here, we report a patient with spinal epidural EMH spanning 11 thoracic vertebral levels, D2 to D12.

CASE REPORT

**Physical presentation and examination**

A 25-year-old male with known thalassemia intermedia, who had never previously required a blood transfusion, presented with a 1-month history of stiffness and weakness in both lower extremities without accompanying sphincteric dysfunction. The general physical examination
revealed palpable splenomegaly, while the neurological assessment demonstrated localized tenderness over the D5 level, with attendant 4/5 weakness in both lower extremities, hyperactive bilateral Patellar and Achilles reflexes with bilateral Babinski responses, and a graded sensory loss to pin appreciation below D5.

**Neurodiagnostic studies**

The magnetic resonance imaging (MRI) of the thoracic spine revealed a posterior, isointense soft tissue epidural mass extending from D2 to D12 on both pulse sequences (T1, T2), suggestive of “red marrow” [Figures 1 and 2]. The cord appeared compressed anteriorly, thinned, and atrophic. The vertebral bodies displayed isointense to hypointense signals on both T1WI and T2WI images along with coarsening of the trabecular pattern; these findings suggested a hypercellular bone marrow consistent with EMH. Of interest, the cervical spine MR was normal.

**Management**

The patient was given multiple blood transfusions and it was decided to repeat a MRI after a period of 8 weeks and further treatment plan will be decided based on the MRI findings.

**DISCUSSION**

**Review of the literature on EMH**

EMH is a compensatory reaction, usually associated with chronic hemolytic anemias. It commonly involves the spleen, liver, lymph nodes, adrenal glands, and pleura. [26,28] Although most patients with EMH are asymptomatic, rare involvement of the spinal epidural compartment may lead to spinal cord compression and myelopathy. Gatto [8] first described this, while other cases studies cited the additional accompanying clinical and MR patterns of progressive lower thoracic spinal cord compression secondary to expansile epidural EMH.[10,13,14,18,27,28] Salehi et al. reported EMH in a 34-year-old male patient with known beta thalassemia. [26] They also reviewed 56 cases described in literature till 2001 and found that out of those 56, 42 cases were due to thalassemia. Only five patients had paraplegia in their study and none of them had thalassemia.

Issargisil et al. [10] studied 12 patients with thalassemia who had spinal cord compression attributed to EMH; these cases involved the lower thoracic region (the commonest location of involvement). Mancuso et al. reported a 15-year-old thalassemia patient presenting with cauda equina syndrome as a result of cauda equina compression.
due to EMH. The incidence of EMH was reported to be 15% in patients with homozygous beta-thalassemia who received regular blood transfusions. In the literature review of Martina et al., male to female ratio was 2.5:1 and most of the affected patients were within 15-45 years of age. The site of compression was the thoracic region in all cases in their series, as seen in our patient.

EMH is diagnosed on the basis of clinical and radiological findings. Symptoms and signs suggestive of compressive myelopathy in a patient with chronic hemolytic anemia should alert one regarding the possible spinal cord compression as a result of EMH.

EMH appears as a lobulated, well defined, and homogenous to heterogeneous soft-tissue mass on computed tomography (CT). Usually there is no evidence of calcification, and bony changes like widening of the medullary cavities of the ribs and coarsening of the trabecular pattern can be appreciated on CT. MRI has now replaced CT as the radiological investigation of choice due to its better soft tissue resolution. EMH appears as a lobular well circumscribed mass and active lesions have an intermediate intensity on both T1- and T2-weighted images and there is no or minimal enhancement following contrast administration. The important differential diagnoses include lymphoma, myeloma, and metastases. The lack of intense contrast enhancement differentiates EMH from other differentials.

Pathogenesis and management

Although the pathogenesis remains controversial and it is proposed that EMH might originate from the stimulation of embryonic rests in the epidural space or it may be a direct extension of the bone marrow. The treatment options include blood transfusion, surgical decompression, agents like hydroxyurea and radiotherapy. Various authors in literature have supported each of these modalities but there is no study comparing each of these modalities due to lack of enough number of cases.

Some authors have used blood transfusion as the sole therapy and have reported improvement of symptoms but improvement is usually incomplete. Blood transfusion acts by down regulating the erythropoietin production. However, it is to be kept in mind that blood transfusion is not completely free of risks and can result in transmission of infections. Patients with mild spinal cord compression can be managed by blood transfusions and more often it is used in combination with other modalities like surgery or radiotherapy. Few authors have also used cytostatic agents like hydroxyurea in combination with transfusion.

Surgical decompression provides immediate decompression of the cord but at the cost of massive bleeding and has been used by various authors. In patients presenting with paraplegia as a result of severe cord compression, surgical decompression may provide benefit to the patient.

Radiotherapy has also been advocated by many authors and has been proposed to halt the overproduction of the overgrown marrow. Malik et al. managed a patient of thalassemia intermedia with paraplegia with radiotherapy alone. There might be initial worsening with radiotherapy due to increased cord edema, and steroids can control the edema. Low doses of radiation usually do suffice due to radiosensitivity of the hematopoietic tissue. Good results with recurrence rates of about 19% have been reported with radiotherapy alone in the literature.

In the literature review by Salehi, 8 patients were managed by surgery alone, while 23 patients with radiotherapy alone. Nineteen patients were managed by a combination of therapies, including various surgery, radiotherapy, blood transfusion, phlebotomy, or steroids.

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