Case Report

CORD COMPRESSION DUE TO EXTRAMEDULLARY HEMATOPOIESIS IN TWO PATIENTS OF \(\beta\)-THALASSEMIA INTERMEDIA

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ABSTRACT

Thalassemia is the most important hemoglobinopathy in Khuzestan province. Thalassemia intermedia (TI) is a genetically heterogenous disease and can result from many different genetic lesions. We report two cases of TI-EMH caused by two separate mechanism and their successful management. Magnetic Resonance Imaging (MRI) is the best diagnostic method in these cases. Management can be done via: Transfusion therapy, Radiotherapy, Hydroxyurea (HU), and Surgery. A 17 years old girl with \(\beta\)-TI previously asymptomatic presented with back pain and leg weakness which started one month ago. The other patient was 25 years old man referred to hospital with back pain, paresthesia, urine frequency and impairment of gait. In the first case the cause of cord compression was the osseous expansion while in the second patient it was related to soft tissue EMH. Next term was successfully treated with low dose radiotherapy and HU. Radiotherapy was initiated with 200cGY fractions to a total dose of 1600cGY and HU 10mg/kg/day. At the end of radiotherapy, the patient was ambulatory with mild residual weakness. She was regularly followed for two years; at present she is active and asymptomatic. The second patient was successfully treated with low-dose radiotherapy and HU. Radiotherapy was started in 200cGY fractions to a total dose of 1600cGY and HU was given at 10mg/kg/day. At the end of radiotherapy the patient was ambulatory with mild residual weakness. He was regularly followed for six months. At his last visit, he was able to walk and climb stair without any assistance. His neurological examination was much better than before. The most common site of spinal epidural extramedullary hematopoiesis is the posterior aspect in the thoracic spine. EMH can be prevented by regular transfusion therapy which corrects anemia and thereby abolishes the stimulus for EMH. Surgical decompression is the method of choice for the management of EMH because histological diagnosis can be established and immediate decompression of the mass can be achieved. This is especially important to decompress the spinal cord quickly in patients with epidural mass caused by EMH. The disadvantages of surgical intervention include risk of excessive bleeding due to high vascularity of the mass and higher incidence of recurrence. In areas where thalassemia is prevalent, EMH should be considered in the differential diagnosis of patients who have chronic anemia with an intrathoracic mediastinal mass.

KEY WORDS: Extramedullary, Hematopoiesis, Thalassemia, Intermedia, Radiotherapy, Hydroxyurea.
thalassemia major. Thalassemia intermedia patients do not receive regular blood transfusions unlike thalassemia major patients. Their bone marrow therefore tries to overcome anemia by increasing its activity. Extramedullary hematopoiesis (EMH) is a rare phenomenon in which apparently normal blood cells are formed out side the bone marrow (primarily affecting the spleen, liver, lymph nodes, spine and thorax).

Thalassemia is common in Iran particularly in south west of Iran. It is estimated that 7-10 percent of population in Khuzestan province is affected by beta thalassemia minor. This syndrome is due to an imbalance of globin chain synthesis. In the case of beta TI, the imbalance is greater than that seen in beta thalassemia trait but less than that in beta thalassemia major. TI has wide range of clinically and genetically heterogeneous patients. Some patients are able to maintain high hemoglobin levels only thanks to huge hemopoietic expansion and bone abnormalities. Masses of heterotopic marrow often develop in patients as a result of continuous erythropoietic stress. EMH is a compensatory mechanism that occurs in patients with hematological dysfunction such as thalassemia major or intermedia and sickle cell anemia. EMH most commonly occurs in organs that have physiological hematopoiesis during embryonic life, especially the liver, spleen and lymph nodes. Other less frequent locations of diffuse compensatory EMH are the kidney, adrenal gland, breasts, spinal cord, plura, pericardium, duramater, adipose tissue and skin. Intra thoracic EMH is a rare condition that is usually asymptomatic and treatment is unnecessary, except in the presence of complications. Massive hemothorax, symptomatic pleural effusion and spinal cord compression are complications of intrathoracic EMH. We report two cases who had regression of intrathoracic EMH with radiotherapy and HU therapy of beta-TI.

History of Patients:
Case-1: A 17-years old girl with beta-TI presented with back pain and leg weakness which started one month ago. In her past medical history, she had blood transfusion for one year between 3 and 4 years with provisional diagnosis of alpha TI. At present she is taking folic acid tablet. On admission, she was not able to climb stairs and run easily.

She had wide based gait. Upper extremities examination was normal, but there was sensory deficit below the level of T5-T6. Deep tendon reflexes were decreased and lower limb strength tone were 3/5, there was no fecal and urinary incontinence. Her liver and spleen were palpable 3 cm below the costal margin. Hemoglobin concentration was 10g/dl, hematocrit: 32.2, reticulocyte count: 6%, mean corpuscular volume: 71.6 fl, mean corpuscular hemoglobin: 21.8 pg, and mean corpuscular hemoglobin concentration: 30.69g/dl. Red blood cell count was 4.49 × 10^{12} cell/liter total billirubin: 4.2 mg/dl, direct billirubin: 0.5 mg/dl serum ferritin: 760ng/ml. Hemoglobin Electrophoresis on the alkaline media of cellulose acetate showed Hb A1:0 Hb F:95% Hb A2:5%. Thoracic spine imaging with and without contrast were performed which revealed that at T2 weighted images demonstrated abnormal bright signal intensity of thoracic vertebrae bodies of T5, T6 and T12 with retro-pulsion which caused thecal sac and cord compression (Fig-1).

Fig-1: Sagittal T2-weighted MR images of the thoracic spine demonstrate bright signal intensity of the vertebrae T5, T6 and T12 which push the cord and make spinal canal lumen focal stenosis. (Arrows)
Cord compression due to extramedullary hematopoiesis

Case 2: A 25 years old man was referred to the hospital with back pain, paresthesia, urine frequency and impairment of gait. He was not able to walk without assistance. Total duration of the disease was 3 weeks. He had no history of thalassemia or blood transfusion. Past medical history included at the age of 9 years old and occasional folic acid supplementation consumption. In general appearance, he had facial expression of pain, seleral icterus, atrophy in lower extremities muscle. On physical examination, his blood pressure was 100/60mmHg, an apical 2/6 systolic murmur was heard, his neurological examination of the upper extremities was normal but in the lower extremities, muscle atrophy was seen. He had ankle clonus, exaggerated deep tendon reflexes, sensory deficit below T5 and T6. Lower muscles strength was 2/5. His hemoglobin concentration was 10.2g/dl , hematocrit: 32.7, reticulo-

cyte count: 4%, mean corpuscular volume:80.5fl, mean corpuscular hemoglobin: 25.1pg, mean corpuscular hemoglobin concentration:31.2g/dl. Red blood cell count:4.06*10¹² cell/liter total billirubin: 3.2mg/dl, direct billirubin:0.5mg/dl and serum ferritin was 1519ng/ml, Hemoglobin Electrophoresis on the alkaline media of cellulose acetate showed Hb A1:0% Hb F:98% Hb A2:2% ,uric acid 8.5 mg/dl,calcium:8.5ms/dl and phosphorous:5mg/dl. AST:74U/L, ALT:81 U/L NRBC:320/100WBC. Chest roentgenogram showed normal size of heart and multiple para vertebral masses due to EMH and coarsening of trabecular pattern of thoracic bone (Fig-3). MRI demonstrated soft tissue density from T2 to T10; causing narrowing of the cord canal (Fig-4).

First case was successfully treated with low-dose radiotherapy and HU. Radiotherapy was started in 200cGY fractions to a total dose of 1600cGY and HU 10mg/kg/day. At the end of radiotherapy, the patient was ambulatory with mild residual weakness. She was regularly followed up for two years, at present she is active and asymptomatic. The second case was successfully treated with low-dose radiotherapy and HU. Radiotherapy was started in 200cGY fractions to a total dose of 1600cGY and HU 10mg/kg/day. At the end of radiotherapy the patient was ambulatory with mild

Fig-2: MRI 43 days later shows complete resolution.

Fig-3: Chest X-ray shows multiple Para vertebra masses, ribs widening, coarse trabecular pattern & osteopenia.

Fig-4: MRI shows long extramedullary mass with intermediated signal on T1 sequence through T2 to T10 and impression on spinal cord.
residual weakness. Forty three days later MRI of the first case showed complete resolution (Fig-2). He was regularly followed up for 6 months. During his last visit, he was able to walk without a stick and he could climb stairs and his neurological examination was much better than before. One month later MRI of the second case showed some resolution and incomplete disappearance of soft tissue masses (Fig-5).

DISCUSSION

Spinal cord compression due to EMH is a well described but rare syndrome encountered in several clinical hematological disorders, including thalassemia, sickle cell anemia, and hereditary spherocytosis.\(^7,9,10,13\) It is rarely seen in Gaucher’s disease, Paget’s disease, alcohol-related macrocytosis and congenital dyserythropoietic anemia.\(^11,14-16\) EMH is secondary to the production of blood cells outside the bone marrow and is a compensatory mechanism for bone marrow dysfunction.\(^6\) EMH is commonly seen at sites such as abdomen, chest, or epidural space.\(^17,18\) There are two forms of EMH namely “para-osseous”- in which the normal medullary tissue of the bone marrow ruptures through the bone to present as a para-osseous mass, and “extra-osseous”- in which EMH occurs within soft tissue. Para-osseous EMH occurs more frequently in hemoglobinopathies whereas extra-osseous EMH accompanies predominantly with the myeloproliferative disorders.\(^19\) Involvement of the epidural space by extramedullary hematopoiesis is most common in patients with thalassemia. It may occur by direct extension from the bone marrow, stimulation of embryonic multipotential hematopoietic stem cells, or via hematogenous emboli. Patients may present with complaints of back pain or spinal cord symptoms. Extramedullary hematopoiesis usually forms a soft, red mass resembling a hematoma on its cut surface. On histologic analysis, all hematopoietic elements are found in extramedullary hematopoiesis.\(^20\) Recognition of spinal cord EMH required prompt physical examination and MRI for accurate diagnosis.\(^21\) Documentation with an imaging technique, such as MRI is mandatory.\(^22\) Bone abnormalities are evident on plain radiographs. Expansion of the bone marrow is seen as osteoporosis with coarsened trabeculation. With severe involvement, there is expansion of the bony cortex. Spinal cord involvement may be suggested by the presence of paraosseous masses seen on chest radiographs. Along with the bone abnormalities, CT demonstrates extramedullary hematopoiesis as a soft-tissue mass that is often adjacent to the involved bone. Extramedullary hematopoiesis can be distinguished from the epidural fat because the former has attenuation similar to that of muscle and shows enhancement with intravenous administration of contrast material.\(^23\) MR imaging is the most effective method of demonstrating extramedullary hematopoiesis in the epidural space and is indicated on an urgent basis when symptoms of spinal cord compression are present. On T1-weighted images, extramedullary hematopoiesis is seen as an extramedullary mass with signal intensity slightly higher than that of the adjacent red marrow of the vertebrae. Similar findings are seen on T2-weighted images, with the signal intensity of extramedullary hematopoiesis being only slightly higher than that of bone marrow. Use of contrast material is unnecessary. T2-weighted sequences may be useful to demonstrate the high signal intensity of an injured spinal cord. This high signal in-
tensity is from edema, myelomalacia, or gliosis of the spinal cord due to chronic compression. The most common site of spinal Epidural extramedullary hematopoiesis is posteriorly in the thoracic spine. Epidural space between the spinal dura mater and the ligaments and periosteum of the posterior spinal elements contains fat, loose connective tissue, and venous plexuses. It is widest in the posterior thoracic spine and is the most common site for extramedullary hematopoiesis when the spinal cord is affected. Management strategies have included radiotherapy, blood transfusion and, occasionally, surgery. Spontaneous recovery with no therapeutic intervention has also been reported but it may take several months to occur and is subject to frequent recurrence and morbidity. Although there is limited experience with HU in thalassemia, some studies have demonstrated successful regression of EMH with HU therapy. The EMH can be managed with radiation, surgery, transfusions, or a combination of these therapies. Radiation in conservative doses of (750-3500 cGy) is non-invasive, avoids surgical risks of potentially severe hemorrhage and incomplete resection and has a high complete remission rate in the majority of patients. Relapse rate moderate (37.5%), but retreatment provides excellent chance for second remission. EMH can be prevented by the institution of regular transfusion therapy which corrects anemia and thereby abolishes the stimulus for EMH. Surgical decompression is the method of choice for the management of the disease because histological diagnosis can be established and immediate decompression of the mass can be achieved. This is especially important in rapid decompression of spinal cord in patients with epidural mass caused by EMH. Disadvantages of surgical intervention include risk of excessive bleeding due to high vascularity of the mass and higher incidence of recurrence. Furthermore, total resection of the mass can lead to clinical decompensation and deterioration since these masses play a crucial role in maintaining an adequate hemoglobin level. To avoid this, incomplete resection of the mass should be attempted, followed by low dose radiation therapy. Being very sensitive to radiation therapy, low doses of radiation (1000-3000cGy), bring about a good clinical response. Major disadvantages of radiation therapy are lack of histological diagnosis and the reduction in the bone marrow activity secondary to the procedure itself. Hydroxyurea, a myelosuppressive agent, has also been successfully employed in the management of EMH. Differential diagnosis of posterior mediastinal location of EMH includes neurogenic tumours, lymphoma, paravertebral abscess and metastatic carcinoma. In geographic areas where thalassemia is prevalent, EMH should be considered in the differential diagnosis of patients who have chronic anemia with an intrathoracic mediastinal mass.

ACKNOWLEDGMENT

Authors wish to thank patients and their families. This work was supported by Physiology and Thalassemia Research Center, Ahwaz Jondishapur University of Medical Sciences, Ahwaz, Iran.

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