E-BETA THALASSEMA WITH EXTRAMEDULAR HEMATOPOIESIS: A CASE REPORT
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ABSTRACT: Extramedullary hematopoiesis (EMH) is a well-recognized process in which the body attempts to maintain erythrogenesis in response to an alteration in the normal production of red blood cells. It is observed in hemoglobinopathies, myeloproliferative disorders, neoplasm's involving the bone marrow and other conditions. Commonly observed areas of EMH include the paraspinal regions of the thorax, liver and spleen, but it has been reported in other locations, including the adrenal gland, bowel, dura mater and breast. Our patient presented with E beta-thalassemia along with hypogonadism who was later diagnosed of having EMH by imaging studies.

KEYWORDS: E-Beta Thalassemia, Extramedulary Hematopoiesis.

INTRODUCTION: Extramedullary hematopoiesis (EMH) is a common compensatory mechanism for chronic anemia found in patients with hemoglobinopathies such as thalassemia, sickle cell anemia, and hereditary spherocytosis. These patients are usually asymptomatic. EMH usually manifests in the thorax as multiple posterior mediastinal or paravertebral masses and masses along the lateral margins of the ribs.¹⁻³ Herein, we report a patient with E beta-thalassemia disease who presented with exertional dyspnea later diagnosed of having EMH by imaging studies along with hypogonadism.

CASE REPORT: A 32 years old male patient presented in the OPD with complaints of generalized weakness for last 2-3months along with dyspnea on exertion for last one month. He also complained of erectile dysfunction and decreased libido for last two years and sparse axillary and pubic hair. He received 30 units of blood for chronic anaemia for last 10 years. There was no history of orthopnea, PND, chest pain or syncopal attack. There was no history of associated fever but he complaints of occasional non-productive cough. No history of any haemoptysis. His bowel and bladder habits are normal.

On examination, patient was emaciated and was pale, icteric and tachypneic. He was tachycardic (110/min) and normotensive (110/70mm Hg). Patient also had testicular atrophy. Systemic examination revealed hepatomegaly and massive splenomegaly. There was grade 4/6 pancontinuous murmur in the mitral area along with loud P2 in pulmonary area. Chest auscultation also revealed bilateral reduced breath sounds and chest resonance was impaired on percussion in bilateral mammary and inframammary area.

Investigations revealed patient was anaemic (Hb- 6.6gm/dl), TLC- 4500/mm³, DLC- N22 L74 M02 E00 B02 and platelet count was 2.4 lakhs/cu mm. Peripheral blood smear showed marked anisopoikilocytosis, polychromasia and hypochromia to severe degree. LFT revealed S. bilirubin- 4.6 mg/dl (Direct-2.6, indirect 2.0), ALP- 162 IU/L, AST- 177IU/L, ALT- 110IU/L, Total protein- 7.4 g/dl, A/G ratio- 1.8:1. Ferritin level was 4090ng/ml. Testosterone level was 157.08ng/ml and free testosterone level was 1.14pg/ml.
Chest X-ray PA View showed bilateral homogenous opacities involving middle zones. CECT thorax also confirmed bilateral well-defined mass with rib erosion. X-ray skull also showed hair on end appearance. USG whole abdomen showed moderate hepatomegaly with huge splenomegaly, extensive splenic micro calcification, dilated PV and SPV and microlith in right kidney. Echocardiography revealed severe MR with PAH. FNAC study of the lung mass revealed it to be a site of extra medullary erythropoiesis.

DISCUSSION: Extramedullary hematopoiesis is seen in a variety of hematologic disorders. It usually develops as a compensatory response in patients with severe haemolytic anaemia such as thalassemia and conditions such as myelofibrosis and myelophthistic anaemia, where there is extensive replacement of normal marrow.

The most common sites of extramedullary hematopoiesis are the liver, spleen and lymph nodes. This condition is rarely manifested as a mass-like lesion within the thorax in patients who have history of frequent blood transfusion and in whom hemoglobin level is kept above 7
g/dL. However, EMH is commonly found in thalassemia-intermedia patients, who usually reach adult life without the need for frequent blood transfusion, because their erythropoiesis is not suppressed.²

EMH masses are usually located in the lower paravertebral areas and are usually multiple and bilateral.¹⁻³

Pathologically they may appear as lobulated dark red-purple fleshy paravertebral masses, without destruction of adjacent ribs and vertebrae, which helps distinguish them from other conditions, such as neurogenic tumors.³ Other sites of intrathoracic EMH include the anterior mediastinum¹ and pleura.⁵

The pathogenesis of intrathoracic EMH includes the extrusion of bone-marrow stem cells through the thin cortex of the vertebral bodies and ribs, abetted by negative pressure; proliferation of the stem cells, which transform into a nodule of hematopoietic tissue upon demand; and proliferation of the embolized hematopoietic tissues from other areas to the intrathoracic region, such as the spleen.¹

CT with intravenous contrast enhancement is a useful imaging method for identifying EMH. The typical CT appearance of EMH is smoothly margined, soft-tissue masses with homogeneous enhancement along the paravertebral regions.⁴,⁶,⁷ The masses may extend into the epidural space and cause spinal-cord compression.⁹,¹⁰ On magnetic resonance imaging, EMH may appear as iso intense paravertebral masses on both T1- and T2-weighted images, with intermediate enhancement after administration of a paramagnetic agent.¹⁰

Technetium⁹ sulfur colloid radionuclide bone marrow scan may show increased tracer activity in the lung and intrathoracic cavity.⁸ Cytological study of the pleural fluid may reveal enlarged mega karyocytes, along with factor-VIII-related antigen immunoreactivity.¹¹ Low-dose radiation should be the treatment of choice, because EMH tissues are highly radiosensitive.⁵

Radiation therapy in conjunction with transfusion can inhibit hematopoiesis and decrease the risk of recurrence.¹² Saxon et al¹³ demonstrated that hydroxyurea can induce regression of EMH and reduce ineffective erythropoiesis in beta-thalassemic patients with bony and spinal complications. Splenectomy may help eliminate the main site of red-blood-cell destruction, as previously reported in patients with hereditary spherocytosis.

REFERENCES:

CASE REPORT


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