Beta-Thalassemia Presenting as an Acute Neurological Complication: A Case Report

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Abstract

Beta-thalassemia is an inherited defect in the rate of synthesis of one or more beta-globin chains of hemoglobin resulting in anemia, hemolysis and ineffective erythropoiesis. Extramedullary hematopoietic tumors may develop in extreme cases of dyserythropoiesis. Acute neurological complications in patients with beta-thalassemia have been reported in association with cerebral ischemia, spinal cord fractures and compression from extramedullary hematopoietic tumors. However extramedullary hematopoietic tumors in the setting of acute blood loss leading to neurological complications have never been reported. We report this case of a 27-year-old woman with beta-thalassemia who presented with acute neurological deficit during immediate postoperative period of cesarean section associated with significant drop in hemoglobin. The Magnetic Resonance Imaging of the thoracic and lumbosacral spine revealed compressive intra-spinal and para-spinal extra-medullary hematopoietic tissue. Patient’s symptoms improved dramatically with conservative management.

Keywords: Beta-thalassemia; Acute neurological complications

Introduction

Thalassemias consist of inherited defects in the rate of synthesis of one or more of the globin chains of hemoglobin [1]. Beta-thalassemia is a condition of impaired production of beta globin chains to varying degree, leading to a relative excess of alpha globin chains, which precipitate within the cell [2]. The resultant anemia, hemolysis and ineffective erythropoiesis form the fundamental pathology for clinical manifestations of beta-thalassemia. In extreme cases of dyserythropoiesis, extra-medullary hematopoietic tumors may develop [3-6]. Acute neurological complications in patients with beta-thalassemia have been reported in association with cerebral ischemia, spinal cord fractures and compression from extra-medullary hematopoietic tumors [7-11]. Other hematological disorders such as leukemias are also reported to have acute neurological complications [12-14]. Review of literature has shown that the neurological complications have been reported with higher frequency in association with thalassemia intermedia [15, 16]. To our knowledge there are no reports of patients in whom hemodynamic shifts during surgery caused a sudden surge in extra-medullary hematopoiesis leading to clinical complications.

Case Report

A 27-year-old African woman from Liberia presented to the hospital with 41 weeks of gestation. The patient was scheduled for an elective cesarean section under epidural anesthesia. The surgery was unremarkable and she delivered a healthy baby. However, four hours into the post-operative period the patient noticed complete loss of sensation and power in the left lower extremity. Physical examination revealed stable vital signs and unremarkable cardiopulmonary system. Abdominal examination revealed massive non-tender hepatosplenomegaly. Evaluation of nervous system revealed normal higher functions and cranial nerves. Sensory system showed loss of sensation in all modalities from L1 to L4 and partial involvement of L5 dermatome. Motor system revealed profound weakness in the left lower extremity, sparing anterior half of the foot. Deep tendon reflexes were normal on right side but left knee reflex was absent with preserved left ankle reflex. There was no saddle anesthesia, bowel or bladder incontinence which implied patchy involvement of spinal nerve roots. Laboratory studies revealed an acute drop in hemoglobin concentration from 10 g/
dl pre-operatively to 7.1 g/dl after the surgery within a span of about 30 hours. The patient underwent a Computerized Tomography (CT) scan of abdomen and pelvis with contrast to evaluate intra abdominal hematoma, which incidentally revealed diffuse coarse trabeculations of the visualized osseous structures, predominantly in the sacrum, with evidence of mild bony destruction and infiltration by soft tissue suggestive of possible extra-medullary hematopoiesis (Fig. 1, 2). The Magnetic Resonance Imaging (MRI) of the thoracic and lumbosacral spine with Gadolinium revealed intraspinal soft tissue deposits from T4 to T9 and L5 to S2 level resulting in significant extra-dural compression. There were also para-spinal nodular soft tissue deposits reported from approximately T7 to T10 and L5 to S2 vertebral levels. The appearance of bone marrow on MRI was suggestive of generalized systemic marrow disorder with a lack of normal fat signal intensity compatible with clinically significant beta-thalassemia (Fig. 3, 4, 5). Packed Red Blood Cell transfusions were recommended, however the patient refused. The patient was started on intravenous Dexamethasone at a bolus dose of 10 mg followed by 4 mg every 6 hours for 3 days. The neurological examination improved dramatically with complete resolution of symptoms upon stabilization of the hemoglobin level.

**Discussion**

We present a young woman with history of thalassemia who presented with acute neurological complications after Cae-sarian section. Differential diagnoses of such presentation include (1) epidural hematoma as a complication of epidural anesthesia causing a secondary compression of the nerve roots, (2) femoral nerve entrapment, (3) vertebral compression fracture and (4) a direct compressive effect on the spinal cord from extra-medullary hematopoietic tissue [3-5, 10, 15, 17, 18]. Imaging studies did not reveal epidural hematoma or a spinal compression fracture. Femoral nerve compression occurs usually secondary to lithotomy position during normal vaginal delivery. The extent of the disease clearly indicates the chronicity of the extra-medullary hematopoietic process. However the acuity of onset puzzled us. We
hypothesize that sudden clinical manifestation of these lesions might have been precipitated by drop in hemoglobin leading to sudden surge in extra-medullary hematopoiesis. The resultant hyperactivity could have further compressed the spinal nerve roots leading to weakness and loss of sensation of left lower extremity. This hypothesis is plausible by rapid improvement in neurological signs upon stabilization of hemoglobin. It might be argued that the administration of steroids might have had a beneficial role. However, the resolution was too rapid to be attributable to steroids alone. It has been shown that hyper-transfusion therapy helps to shrink extra-medullary hematopoietic tumors and alleviate the compressive effect on spinal cord [3, 5, 6, 9].

In conclusion, acute neurological complications from extra-medullary hematopoietic tissue are rare but serious complications in patients with extreme dyserythropoiesis as a result of beta-thalassemia that might require blood transfusions and high dose steroids. Acute blood loss may lead to surge in extra-medullary hematopoiesis leading to worsening of compressive effects. Therefore in patients with thalassemia undergoing surgery where significant blood loss is anticipated, prophylactic blood transfusions and close monitoring may be necessary to avoid and manage acute neurological complications [4, 6, 9, 18-20].

References

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