Atypical Presentation of a Case of Hemoglobin E Trait with Gaucher’s Disease

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Abstract: Hemoglobin E heterozygotes (Hb AE) are asymptomatic and homozygotes (Hb EE) have a mild microcytic anemia [1]. However, we had a 2 year old female child presenting with moderate pallor necessitating blood transfusions at 6 months to 1 year interval starting from eight months of age. Thorough clinical examination and investigative work-up revealed Hb E trait with Gaucher’s disease. To the best of our knowledge, we are reporting for the first time the association of Hb E trait with enzyme-study confirmed Gaucher’s disease. Earlier, there have been three case reports of Gaucher-like cells or Pseudo-Gaucher cells (not Gaucher’s disease) associated with classic beta thalassemia major, Hb E disease (HbEE) and thalassemia intermedia [2][3]. It is therefore desirable to understand whether the co-existence of a thalassemia-like syndrome and a storage disease like Gaucher’s disease could influence the clinical expression of each other.

Keywords - Bone marrow biopsy, Gaucher cells, HbE trait, Liver biopsy, Leukocyte betaglucosidase, plasma chitotriosidase

Abbreviations: dl : deciliter, Fig. : Figure, Gm : Gram, Hb : Hemoglobin, HBsAg : Hepatitis B surface antigen, HCV : Hepatitis C virus, HPLC : High-performance liquid chromatography, mm³ : cubic millimeter, RBC : red blood cells, RDW : Red cell distribution width, PAS : Periodic acid-Schiff, Thal : Thalassemia

I. INTRODUCTION

HbE trait is an asymptomatic entity on its own, with no clinical significance, unlike compound heterozygous state (HbE- Thal). However the need for repeated blood transfusions in Hb E trait (Hemoglobin electrophoresis and HPLC proven) especially associated with moderate splenomegaly called for further work-up and confirmed Gaucher’s disease, thereby explaining the unusual requirement of several blood transfusions in Hb E trait. We strongly believe that this is the first such report of a case of Hb E trait with Gaucher’s disease.

II. CASE REPORT

A 2 ¹⁄₂ year old girl presented to us with moderate pallor and abdominal enlargement. On palpation, the liver was enlarged 5 cm below the costal margin with left lobe enlarged 6 cm at the epigastrium. It was firm in consistency with a well defined margin. The spleen was palpable about 10 cm along its long axis. She had her first blood transfusion at 8 months of age at a hemoglobin level of 6.7 gm/dl with some evidence of hemolysis in the form of hypochromia, microcytosis, anisopokilocytosis and polychromasia. Her platelet count started showing a rapid decline over the next 6 months (as suggested by her sequential reports) though the haemoglobin level was maintained in the range of 7-11 gm/dl. The leukocyte count showed a gradual falling trend too. Hb electrophoresis of the mother (in alkaline buffer pH 8.6) showed HbE (by elution) to be 28.6% and Hb F (by Singer’s method) as 0.6%. The electrophoresis pattern showed a major band at AF region and a less prominent level was maintained in the range of 7-11 gm/dl. The leukocyte count showed a gradual falling trend too. Hb electrophoresis of the mother (in alkaline buffer pH 8.6) showed HbE (by elution) to be 28.6% and Hb F (by Singer’s method) as 0.6%. The electrophoresis pattern showed a major band at AF region and a less prominent band at E region giving the impression of heterozygous HbE trait. The father showed a normal Hb electrophoresis pattern. An earlier HPLC report of our patient at one and half years of age showed Hb F to be 2%, Hb A₂ +E to be 28%, the impression being given of HbE trait.

As we could not explain why HbE trait would cause so much anemia or splenomegaly when she presented to us for the first time at two and half years of age, we carried out a thorough laboratory work-up.

A blood count revealed Hb level of 6.9 gm/dl, and a reduced platelet count of 7000/ cu mm. The peripheral smear showed normocytic and microcytic hypochromic RBCs with moderate thrombocytopenia. Nucleated RBCs were absent.

Bone marrow aspirate (Fig. 1) revealed storage cells with basophilic fibrillar cytoplasm, eccentric nucleus resembling Gaucher’s cells present in good numbers. There was normal maturation of megakaryocytic, granulocytic and erythroid cells.

Bone marrow trephine biopsy (Fig. 2) revealed sheets of large cells replacing hematopoietic marrow over many areas. These cells showed single or multiple small irregular nuclei with fine nuclear chromatin and abundant cytoplasm. Many of the cells showed wrinkled paper appearance of the cytoplasm. Islands of...
hematopoietic marrow with sequential maturation were also present. Reticulin stain showed intersecting fine reticular fibers over focal areas. All these findings were suggestive of Gaucher’s disease.

Serum ferritin was marginally increased. Tests for HBsAg and anti-HCV antibody were non-reactive. Serial blood counts showed Hb in the range of 7-9 gm/dl, a rapidly declining platelet count with a minimum recorded count of 52000/mm³ and a falling trend in leukocyte count (minimum = 3800/mm³). Hypochromic, microcytic anemia with normal Red cell distribution width (RDW) were consistently found.

Prothrombin time and Activated Partial Thromboplastin time were within normal limits. Liver biopsy showed focal collections of macrophages having abundant, pale, acidophilic cytoplasm with some showing striations. These cells showed diastase resistant PAS positive material. All these findings were strongly suggestive of Gaucher’s disease. Finally, Enzyme analysis confirmed the diagnosis of Gaucher’s disease. Beta glucosidase in leukocytes was markedly reduced and in plasma chitotriosidase was significantly increased, thereby establishing the diagnosis of Gaucher’s disease Type I.

III. Discussion

Hb E trait, the second most common globin mutation worldwide, is asymptomatic in heterozygotes and causes mild microcytic anemia in homozygotes¹. Our case, about 2½ years old on first presentation had four units of packed RBCs since her first transfusion at eight months of age. The child maintained a Hb level of 7-11 gm/dl in between transfusions. Though the frequency of transfusions had not been high, the need for blood transfusion in haemoglobin E trait was rather unusual. The child had a normal facies and a firm splenomegaly and hepatomegaly with increments in splenic size of 4 cm and in liver size of 2 cm in the two and half year of her follow up with us. As of now, the spleen is palpable 14 cm in its long axis and the liver is palpable 7 cm in the midclavicular line and 8 cm at the epigastrium. A rapid decline in the platelet count and gradual decline in leukocyte count could be explained by the hypersplenism resulting from massive splenomegaly and also the bone marrow infiltration by Gaucher’s cells. The moderate degree of microcytic hypochromic anemia necessitating intermittent blood transfusions can be explained by the chronicity of the disease state and with further aggravation by nutritional imbalances. The normal reticulocyte count and RDW have negated the possibility of any haemolytic factor in the causation of anemia. The absence of evidence of any hemophagocytosis by Gaucher’s cells in the histopathology of the bone marrow biopsy specimen and only a slightly raised serum ferritin further ruled out hemolysis as a major cause of anemia.

Gaucher-like or PseudoGaucher cells have been reported by a few authors in association with classical beta thalassemia major, Hb E disease (EE), thalassemia intermedia, chronic myeloid leukemia, multiple myeloma, etc. However, either enzyme study for beta glucosidase in leukocytes or skin fibroblasts or a search for birefringent crystals in Gaucher like cells have ruled out Gaucher disease. All these conditions are characterized by enhanced cell turnover with accumulation of beta-glucosidase enzyme in the lysosomes of the mononuclear phagocyte cells by the cell membrane derived glycosylceramide ². These represent a relative rather than absolute deficiency of the enzymes in the histiocytes ³⁴. Interestingly in one report of monoclonal gammopathy of undetermined significance, these cells had the morphological and immunological features of plasma cells rather than macrophages⁶. However other authors concluded on immunoelectrophoresis and electron microscopy, that the Pseudo-Gaucher cells even in a case of myeloma were marrow macrophages engorged with immunoglobulin ⁷. While most described these cells in the bone marrow, a few reports document them at other locations like peritoneal fluid ⁸, spleen ⁹¹⁰ and lymphnode ¹¹¹². In our patient the presence of sheets of Gaucher cells in the bone marrow biopsy specimen, focal collection of PAS positive & diastase resistant Gaucher cells in liver biopsy specimen, a reduced beta glucosidase activity in leukocytes and a significant increase in plasma chitotriosidase¹³ clinched the diagnosis of Gaucher disease Type I in association with HbE trait.

IV. Conclusion

This report stresses the need to always look for the presence of storage disorders like Gaucher’s Disease in any clinical case presenting as thalassemia –like syndrome where the hemoglobin electrophoresis report does not match with the severity of symptoms. Though the presence of Gaucher like cells has been reported by a few other authors in association with thalassemia –like syndromes, Gaucher’s disease was never documented in their reports. Our case, we believe, is probably the first such report. There is also a need to evaluate whether the co-existence of a thalassemia-like syndrome with a storage disorder might influence the clinical course of each other.
REFERENCE.


Fig. 1: Bone marrow aspirate revealed storage cells with basophilic fibrillar cytoplasm, eccentric nucleus resembling Gaucher’s cells (High Power)

Fig. 2: Bone marrow trephine biopsy revealed sheets of large cells replacing hematopoietic marrow over many areas (High Power)