Case Report

Type-1 Glanzmann’s thrombasthenia: a rare cause of epistaxis in a child

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ABSTRACT

Glanzmann’s thrombasthenia (GT) is a rare genetic platelet surface disorder of glycoprotein IIb/IIa receptor presenting with mucocutaneous bleeding of varying severity. We are reporting an unusual case of a child presenting with recurrent epistaxis with prolonged bleeding time, moderate thrombocytopenia and giant platelet size. GT was suspected because the platelet aggregation was abnormal with adenosine diphosphate, epinephrine, collagen, and thrombin; but normal with ristocetin. Diagnosis was confirmed by flow cytometry which showed deficiency of platelet membrane receptors CD 41 (Gp IIb) and CD 61 (GpIIIa) with normal expression of CD 42b (GpIb). Platelets transfusions and antifibrinolytics were given to manage bleeding. Due to repeat platelets transfusions patients with GT can develop anti-platelet antibodies for which rFVIIa (recombinant activated factor VII) is effective. Definitive treatment includes stem cell transplant or gene therapy.

Keywords: Epistaxis, Glanzmann’s thrombasthenia, Platelet transfusion

INTRODUCTION

Glanzmann’s Thrombasthenia (GT) is a rare autosomal recessive platelet disorder. It was first described by Dr. Edward Glanzmann in 1918.1 Reported incidence of GT is one in million.2 A higher incidence is seen in families with consanguineous marriages, with equal sex predilection.3 Patients with GT present with mild to severe mucocutaneous bleeding manifestations.3 GT is usually characterised by normal platelet number and morphology, although, rarely, moderate thrombocytopenia and giant platelet size are also observed.4,5 The platelet aggregation is normal with ristocetin but abnormal with physiologic agonists including adenosine diphosphate (ADP), epinephrine, collagen, and thrombin.5

Here, we are reporting a case of GT with unusual features of thrombocytopenia and giant size platelets who presented with epistaxis.

CASE REPORT

A 4 years old Muslim male child, born of a consanguineous marriage, 2nd by birth order, resident of Kutch district; presented to pediatrics outpatient department (OPD) with complaint of recurrent unprovoked epistaxis on and off for two years. On physical examination, severe pallor was present and rest of the examination was normal. Patient had history of similar episodes of epistaxis in the past for which blood was transfused on two occasions. The initial blood investigations showed microcytic hypochromic anemia (hemoglobin i.e. Hb 5 gm %), normal leucocyte count (8000/ml), reticulocyte count (0.6 %), erythrocyte sedimentation rate (ESR) (12 mm/hour), and C-reactive protein (CRP) (<1 mg/dl); and, decreased platelet count (1.2 lakhs per microliter), serum iron (40 mcg/dl), vitamin B12 (150 pg/ml) and folic acid (4 ng/ml) levels. So, the child was transfused 2 units of packed cell volume. Otorhinolaryngology reference was done for epistaxis and
anterior nasal packing was done. Though the severity of bleeding decreased, steady oozing continued along the edges of nasal packs and also in nasopharynx. After 36 hours, once the bleeding stopped, nasal packs were removed and nasal examination done. Nasal endoscopy showed widespread raw nasal mucosa and no mass lesion or focal bleeders. Computed tomographic (CT) scan of nose and para nasal sinuses did not show any abnormality.

The above findings prompted us to search for possible hematological disorders as cause of recurrent epistaxis. A repeat peripheral blood smear examination showed decreased platelet counts (80,000 per microliter) and giant platelets (Figure 1). However, the bleeding time was 13 minutes (prolonged), favoring the diagnosis of functional disorder of platelets. The child again had an episode of epistaxis for which platelet transfusion was done, oral tranexamic acid, iron, vitamin B12 and folic acid were given. Bleeding stopped, thereby, affirming the probable diagnosis of functional disorder of platelets. Ristocetin induced platelets aggregation test was normal but platelet aggregation with ADP, epinephrine and collagen was not seen. Finally, flow cytometry platelet analysis was done which showed deficiency of platelet membrane CD 41 (Gp IIb) and CD 61 (GpIIa) with normal expression of CD 42b (GpIb) (Figure 2). As expressions of CD 41 and CD 61 were negative, diagnosis of type-I GT was made.

DISCUSSION

GT is a rare autosomal recessive bleeding disorder characterised by defect in platelet aggregation. It is caused by either quantitative or qualitative defect of platelet membrane glycoprotein IIb/IIIa (integrin αIIbβ3) which is essential for platelet aggregation. It is due to genetic defect on ITGA2B or ITGB3 gene on chromosome 17 (12q21). According to integrin αIIbβ3 (CD41/CD61) concentration and its functionality, there are three types of GT. In type I and II, GP IIb/IIIa is either absent or < 5% and 5-20 % of normal respectively. In type III or variant, GP IIb/IIIa is >20% but functionally impaired. Classical GT is characterized by normal platelet numbers, normal platelet morphology, prolonged bleeding time (BT), abnormal clot retraction and absent or decreased platelet aggregation to physiologic agonists like adenosine diphosphate (ADP), epinephrine, collagen, and thrombin but normal platelet aggregation with ristocetin. In our case thrombocytopenia and giant size platelets were present which is unusual of GT but has been reported in some patients. In our case, since platelet aggregation was normal with ristocetin, Bernard-Soulier syndrome, which is another platelet function disorder, was excluded.

Severity of bleeding varies in patients with GT. Some patients have mild muco-cutaneous bleeding while others can have life threatening hemorrhage. Common bleeding manifestations are epistaxis (73%), gingival hemorrhage (55%), purpura (86%), menorrhagia (98%); rarely gastrointestinal hemorrhage (12%), hematuria (6%), hemarthrosis (3%), intracranial hemorrhage (2%) and visceral hematoma (1%) are seen. In our case recurrent epistaxis was main manifestation.

Diagnosis of GT is confirmed by flow cytometry in which platelet membrane CD 41 (Gp IIb) and CD 61 (GpIIa) are decreased with normal expression of CD 42b (GpIb). In our case, CD 41 and CD 61 expressions were negative in flow cytometry so diagnosis of type-1 GT was made. Molecular genetic study can be done to know the site of mutation. Treatment of GT requires platelets transfusion, antifibrinolytics or recombinant activated factor VII (rVIIa) alone or in combination. Definitive treatment by stem cell transplant and gene therapy is required in severe...
cases. In our case, platelet transfusion and antifibrinolytics were given. Generally, in GT the incidence of severe bleeding decreases with age. During follow up of our patient, two episodes of nasal bleeding occurred during next one year for which antifibrinolytics and platelets transfusion were given.

**CONCLUSION**

GT is a rare bleeding disorder, which should be considered as a differential diagnosis in patients with recurrent bleeding manifestations with normal coagulation profile. Rarely, thrombocytopenia or giant platelets can be seen in GT.

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