Fulminant Essential Thrombocythemia Associated with Acquired Von Willebrand Syndrome and Bleeding Episodes in a 14-year-old Girl

C. Schneider¹  E. Stutz-Grunder¹  S. Lüer¹  P. Keller²  J. A. Kremer Hovinga²  R. A. Ammann¹  A. Karow¹

¹Division of Pediatric Hematology/Oncology, Department of Pediatrics, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland
²Department of Hematology, University Hospital/Inselspital Bern and University of Bern, Bern, Switzerland

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Address for correspondence  C. Schneider, Dr. med., Division of Pediatric Hematology/Oncology, Department of Pediatrics, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland (e-mail: christine.schneider@insel.ch).

Introduction

Essential thrombocythemia (ET) is a chronic myeloproliferative neoplasm (MPN) with clonal megakaryocyte proliferation. It is an extremely rare cause of thrombocytosis in childhood and adolescence with an estimated annual incidence of only 0.09 per million. Usually, clinical symptoms associated with essential thrombocythemia are mild or absent.

Background  Essential thrombocythemia is a chronic myeloproliferative neoplasm. It is extremely rare in children below 15 years of age with an estimated annual incidence of only 0.09 per million. Usually, clinical symptoms associated with essential thrombocythemia are mild or absent.

Case  Here, we present the case of a 14-year-old female patient fulminantly presenting with acute symptoms comprising visual impairment, palmar and plantar stabbing pain. Blood count revealed massive thrombocytosis of 2373 × 10⁹/L. Bone marrow morphology showed elevated numbers of mature megakaryocytes. Von Willebrand factor activity/antigen ratio was significantly reduced compatible with an acquired Von Willebrand syndrome associated with high platelet counts. Molecular analyses for driver mutations of myeloproliferative neoplasms including JAK2V617F, CALR and MPL were negative. Acute therapy comprising hyperhydration and oxygen supply complemented by acetylsalicylic acid led to amelioration of symptoms. Medication with hydroxycarbamide maintained a significant reduction of platelet counts but had to be reduced or withheld several times due to neutropenia. Repeated bleeding episodes observed in the course were clearly associated with increases in platelet counts above 1200 × 10⁹/L explained by acquired von Willebrand syndrome. Sixteen months after diagnosis, therapy was switched to pegylated interferon and platelet counts could be stabilized without significant side effects.

Keywords
► essential thrombocythemia
► myeloproliferative neoplasm
► Von Willebrand syndrome

Abstract

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somatic mutations are rarely present in childhood ET. Therefore, diagnosis of childhood ET is challenging and World Health Organization (WHO) diagnostic criteria for adult MPN cannot be simply adopted. In general, ET is associated with a good prognosis, but life expectancy is limited when thromboembolic complications or progression to post-ET-myelofibrosis or acute myeloid leukaemia (AML) occurs. Here, we report on a case of a 14-year-old girl with an impressively fulminant clinical presentation of ET. Symptoms were highly acute and comprised headache, visual impairment and painful paresthesia of the upper and lower extremities. Extraordinarily high platelet counts were found and urgent intervention was required.

**Case Report**

A 14-year-old female patient consulted her general practitioner with recurrent stabbing pain and palmpoplantar erythema since 2 weeks. Blood count performed to exclude underlying infection revealed massive thrombocytosis above the diagnostic range of the analyser used and the girl was referred to the emergency department of our university hospital. Two weeks before, when the patient consulted her general practitioner because of first mild pain episodes, blood count had been performed for the same reason and showed platelets of $1310 \times 10^9/L$. Retrospectively, elevated platelet counts of more than $600 \times 10^9/L$ for the past 3 years and $996 \times 10^9/L$ 12 months before diagnosis had been documented, measured in blood counts examined for suspicion of iron- and vitamin D deficiency and fever. These results had not led to further investigations.

Further history revealed paresthesia of both arms and intermittent headache with impaired vision. Vital signs and clinical findings, including a detailed neurological examination, were normal. In particular, there were no signs of secondary causes for thrombocytosis, i.e. infection, trauma or recent surgery. Apart from vitamin D substitution there was no other medication nor any drug abuse. The adolescent had so far been healthy. The patient’s parents were not consanguineous and had immigrated from Afghanistan when the patient was 9 years old. Apart from multiple miscarriages on the father’s side of the family, family history was unremarkable with no thromboembolic events or bleeding complications. Platelet counts of all first-degree relatives were below $300 \times 10^9/L$.

At our institution, the complete blood count was repeated and confirmed thrombocytosis of $2373 \times 10^9/L$ determined by a Sysmex XN 9000 platelet counter. Haemoglobin and a differential leukocyte count were within age-specific range. Laboratory testing showed a level of C-reactive protein $<3$ mg/L. Serologic testing for Epstein–Barr virus, cytomegalovirus, hepatitis A, B, C and human immunodeficiency virus was negative. Further diagnostics showed a prolonged partial thromboplastin time of 45 seconds (normal range: 25–36 seconds). Because of the known relationship of very high platelet counts ($>1000 \times 10^9/L$) and acquired von Willebrand syndrome, corresponding laboratory analysis was performed as an explanation for the prolonged partial thromboplastin time. Other reasons for the prolonged partial thromboplastin time were not found. No use of oral anticoagulants or heparin was reported. Factor VIII plasma activity (95%) was measured later in the course. Lupus anticoagulant and anti-cardiolipin antibody titre were not determined.

Both von Willebrand factor (VWF) activity (27%) and VWF antigen (48%) were reduced with a pathological VWF activity/antigen ratio of 0.56 (normal $\geq 0.7$). Multimer analysis of VWF revealed a loss of the largest and relative reduction of large VWF multimers as well as accentuated satellite bands (Fig. 1, lane 1).

Ultrasound showed slight splenomegaly. Bone marrow histology demonstrated increased numbers of mature megakaryocytes exhibiting “staghorn” nuclei. In addition, mild fibrosis without typical features of primary myelofibrosis (increased reticulin or collagen fibres, hyperchromatic or ‘cloud-like’ nuclei) was seen (Fig. 2).

Based on these classical findings, we diagnosed ET. However, histologically, there is no clear distinction between a congenital thrombocytosis and ET and also a ‘prefibrotic’ primary myelofibrosis could not be entirely excluded.

Screening for the most common somatic mutations known to be associated with MPN using clinical routine assays was negative for JAK2V617F, MPL W515L/K and deletions in calreticulin, respectively. Translocation of BCR-ABL1 was also ruled out. In addition, we analysed the mutational

**Fig. 1** Von Willebrand factor (VWF) multimer analyses. Separation of von Willebrand factor multimers in a medium resolution (1.7% agarose) gel. Lanes N: normal human plasma pool; patient’s samples (lanes 1–4); lane 1: at diagnosis (platelet count: $2505 \times 10^9/L$); VWF ratio: 0.45; the loss of the large VWF multimers is clearly visible, satellite bands are slightly more prominent than in normal human plasma pool; lane 2: 2 months later (platelet count: $1243 \times 10^9/L$; VWF ratio: 0.65); lane 3: 10 months after diagnosis (platelet count: $688 \times 10^9/L$; VWF ratio: 0.79); lane 4: 11 months after diagnosis (platelet count: $267 \times 10^9/L$; VWF ratio: 0.88); lane 5: plasma of a patient with genetically confirmed Von Willebrand disease 2A (IIA) (VWF ratio: 0.25) characterized by loss of large and medium sized VWF multimers and clearly accentuated satellite bands.
spectrum in our patient using a targeted next-generation sequencing panel comprising all exons and the flanking regions of JAK2, MPL and TPO and more than 60 cancer-related genes. However, even this approach did not detect any clonal aberration.

Because of massive thrombocytosis with associated neurological symptoms, treatment with hydration (2–3 L daily) and oxygen supply was initiated and oral acetylsalicylic acid (100 mg increased to 200 mg divided in two daily doses) was started. Cerebral ischaemia was excluded by an emergency magnetic resonance imaging (MRI) scan.

A therapy with oral 20 mg/kg hydroxyurea (Litalir) daily was started and gradually increased to 40 mg/kg daily. Due to persisting symptomatic thrombocytosis 2 weeks after initiation of treatment, platelet apheresis was performed. Since the patient’s initial ferritin of 34 µg (range: 7–140 µg) was always in a low normal range with borderline haemoglobin levels of 120 g/L (range: 121–154 g/L), we decided to repeatedly apply iron sucrose to minimize any possible additional reactive thrombocytosis due to effective iron deficiency. However, we observed no effect on platelet counts.

Because of neutropenia, hydroxyurea had to be reduced twice and stopped three times for several days, respectively. As a consequence, platelet counts repeatedly increased significantly at around 3 weeks after stopping hydroxyurea (Fig. 3).

Due to these fluctuations of cell counts and the convincing experience with pegylated interferon in adult patients, we replaced hydroxyurea 16 months after the first presentation and switched to treatment with peginterferon alfa-2a (Pegasys) as a subcutaneous injection starting with 90 mcg/week and continuous increase of the dose up to 180 mcg/week over 3 months.

Under this treatment, the patient remained asymptomatic, platelet counts appeared stabilized at around 800 × 10⁹/L and no side effects were observed. Persistent reduction of the platelet count also led to normalization of the VWF ratio.

Notably, there were three episodes with increased bleeding tendency in the further course, two with epistaxis and one with significantly prolonged menstrual bleeding. All of these episodes were associated with an increase in platelet counts (Fig. 3).

Follow-up bone marrow biopsy taken 1 year after diagnosis showed stable histologic features.

Fig. 3 illustrates the clinical course and platelet count over time with respect to different therapeutic interventions.
Discussion

The most common cause of elevated platelet counts in adolescence is reactive, mostly infection-associated, thrombocytosis. These cases are self-limiting and remain asymptomatic. Complications like bleeding or thrombosis are hardly seen in reactive thrombocytosis.6,7

Other reasons for reactive thrombocytosis are surgery, anaemia in particular due to iron deficiency, drug side effects, allergy,7,8 chronic inflammatory disease and asplenia.9

One further but rare differential diagnosis is congenital thrombocytosis, an inherited autosomal-dominant, autosomal-recessive or X-linked condition. However, in our case all family members of the first generation had platelet counts below 300 × 10^9/L and previous blood samples from our patient taken by the family physician showed lower platelet values gradually increasing over months. These findings and on the absence of mutations in TPO and MPL as shown in our diagnostic screening rendered the diagnosis of congenital thrombocytosis unlikely and we diagnosed ET.

ET is an extremely rare cause of thrombocytosis in adolescence. According to the current WHO criteria, diagnosis of ET is made when the following criteria4 are fulfilled: (1) prolonged platelet count over 450 × 10^9/L, (2) bone marrow aspiration showing an elevated number of mature megakaryocytes, (3) not matching one of the WHO criteria for other chronic Philadelphia-negative myeloid diseases and (4) confirmation of one acquired somatic mutation or clonal marker. If the last criterion is not met, the diagnosis of ET can be made after 1 year without evidence of reactive thrombocytosis. In our patient, no somatic mutation or clonal marker was identified while all other criteria were present. Gene mutations typically found in adult MPN are often lacking in children and adolescents suggesting a different disease pathology.10

Only very few paediatric patients with ET present with microcirculatory symptoms, thromboembolic events, haemorrhagic diathesis or splenomegaly. Symptoms in our patient were uncommonly severe and together with extremely high platelet numbers prompted immediate initiation of treatment.

Patients with ET have an elevated risk of thrombotic events but also may show a bleeding tendency. The latter is explained by the quantitative and qualitative platelet alterations resulting in thrombocytopenia which is, as in our case, often aggravated by an acquired Von Willebrand syndrome, the result of adsorption of large VWF multimers on platelets.11–13 Platelets with increased degradation might explain the overproportional reduction of VWF activity with a decreased activity/antigen ratio. These mechanisms have likely caused the episodes of increased bleeding tendency with epistaxis and prolonged menstrual bleeding observed in our patient. It cannot be ruled out that the medication with aspirin augmented this bleeding tendency. The initial therapeutic decision to apply aspirin, however, was based on the significantly increased risk of thromboembolic complications associated with the extremely high platelet counts found in our patient with ET. The two bleeding episodes observed in our patient were clinically not severe. After these minor bleeding episodes, we intermittently halved the aspirin dosage. Since bleeding symptoms occurred in phases of increased platelet counts >1200 × 10^9/L and subsided when platelet counts dropped, it can be assumed that they were mainly caused by the secondary Von Willebrand syndrome.

Regarding the low numbers of treated children with ET and the long-term implications in this age group, the therapeutic approach of childhood and adolescence MPN remains challenging.

We have chosen to use hydroxyurea as primary cytoreductive therapy in our patient because of our experience with this medication in sickle cell anaemia patients where hydroxyurea is used for the induction of foetal haemoglobin and acts as an endothelial cell adhesion modulator.14 However, an aggravated leukaemogenic potential in adult ET patients after long-term use is still controversially debated. In addition, due to its potential fetotoxic and embryotoxic effects, hydroxyurea will have to be replaced for pregnancy.

Alternative therapeutic options are anagrelide and pegylated interferon (Pegasys). Both substances are established in the treatment of adult MPN.15

Like polycythemia vera, ET can transform to myelofibrosis or AML. No prognostic markers are known to predict transformation. Due to the mild bone marrow fibrosis observed in our patient however, we could not entirely exclude ‘pre-blastic’ primary myelofibrosis which is associated with a less optimal prognosis.

In summary, diagnosis of ET is very rare in children and adolescents and should be considered as a differential diagnosis in persistent thrombocytosis only after exclusion of secondary causes. Childhood ET appears to have a different genetic background than adult ET and often mutational status cannot serve as a diagnostic criterion. Paediatric patients with ET are mostly asymptomatic and only a very small minority present with microcirculatory symptoms, thromboembolic events, haemorrhagic diathesis or splenomegaly. Lacking a standard therapy in children and adolescents, management and especially long-term treatment of ET in these age groups can be challenging.

Conflict of Interest
The authors have no conflict of interest to disclose.

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