Bortezomib induces long term remission in children with immune thrombotic thrombocytopenic purpura, refractory to plasma exchange, glucocorticoids, and rituximab: a report on two cases

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Abstract

Acquired thrombotic thrombocytopenic purpura (TTP) in children is a rare but severe disease, which is caused by Immunoglobulin G antibodies, which inactivate a disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13 (ADAMTS13). Daily high-volume plasma exchange (PEX) and immunosuppression with glucocorticoids and rituximab is the current standard of treatment for TTP. We report two females aged 5 and 12 years, with TTP, induced by anti-ADAMTS13 inhibitory antibodies who relapsed very shortly after PEX, rituximab and glucocorticoids, in whom long-term remission with disappearance of ADAMTS13 inhibitors was achieved after one course of bortezomib.

Introduction

Acquired thrombotic thrombocytopenic purpura (TTP) in children is a rare but severe disease, which can lead to devastating critical organ damage or even death.1 It is caused by Immunoglobulin G (IgG) antibodies, which inactivate a disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13 (ADAMTS13), which cleaves the ultralarge von Willebrand factor (ULVWF).2 In the absence of active ADAMTS13, ULVWF interacts with platelets, inducing the disseminated formation of platelet-rich thrombi both in circulation and in microvessels, which can result in severe ischaemic tissue damage.

Daily high-volume plasma exchange (PEX) and immunosuppression with glucocorticoids and rituximab is the current standard of treatment for TTP, which has decreased mortality rates to less than 10%.4-7 Recently, caplacizumab, a nanobody which blocks the A1 domain on ULVWF and prevents its interaction with platelets, has been shown to shorten the interval to remission and also to reduce the likelihood of early death.8 However, a significant proportion of patients have TTP refractory to PEX and immunosuppression, or relapse after achieving remission. We describe the cases of two children with immune-mediated TTP, who relapsed shortly after treatment with PEX, glucocorticoids, and rituximab, and subsequently went into continuous complete remission after one course of bortezomib.
**Patient 1**

A 12-year-old girl of Hispanic origin, presented with petechial rash and bruising, jaundice, backaches, nausea, and arthralgia. A complete blood count (CBC) showed thrombocytopenia (11 x 10^9/L), anaemia (Hb: 9.8 g/dl), reticulocytosis (16.9%), and normal WBC (5.9 x 10^9/L). The total bilirubin level was 48 μmol/L, lactate dehydrogenase (LDH) was 514 U/L, and the Coombs test was negative. A diagnosis of immune thrombocytopenia (ITP) was made, and intravenous immunoglobulin (IVIG) 1 g/kg body weight (bw) was administered, with no resulting increase in platelets. Subsequently, TTP was suspected, and testing revealed severely decreased activity ADAMTS13, <5%, which confirmed the diagnosis of TTP and the patient was transferred to our centre. Upon admission, her examination was remarkable for fatigue, cutaneous haemorrhages, and uterine bleeding. The patient’s Hb was 5.9 g/dl, schistocytes were 16%, reticulocytes were 51%, platelets were 16 x 10^9/L, and total bilirubin was 54.5 μmol/L. Daily PEX was initiated concomitant with methylprednisolone 1000 mg for three days i.v. and rituximab 375 mg/m^2 twice weekly for three weeks. After 18 PEXs, the patient’s platelet count stabilised above 150 x 10^9/L. However, ADAMTS13 was persistently <5%, and a high level of the ADAMTS13 factor inhibitor persisted. Two days after PEX was electively stopped, the patient’s platelet counts dropped again, triggering the resumption of PEX twice weekly, which maintained the platelet count above 300 x 10^9/L. Six weeks after the patient’s 29th PEX, TTP occurred again, with a drop of platelets to 10 x 10^9/L. Thus, because the patient developed TTP refractory to PEX, rituximab, and glucocorticoids, we decided to start therapy with bortezomib, 1.3 mg/m^2 on days 1, 4, 8, and 11. After two additional PEXs and one course of bortezomib the patient’s platelet counts, Hb, and LDH all recovered. ADAMTS13 activity reached 81%, and remained within normal limits during 5 years of follow-up. She received one additional course of bortezomib with 21 days interval. There was no toxicity from bortezomib observed, and IgG serum levels were checked several times during follow-up period, during which they never dropped below 7 g/L, and at the last follow-up was 12.4 g/L (figure 1).

**Patient 2**

A 5-year old girl was admitted to a hospital with petechiae and ecchymoses after a mild respiratory illness. CBC showed thrombocytopenia (5 x 10^9/L), low Hb (6.8 g/dL), and normal WBC (5.5 x 10^9/L). The patient was diagnosed with ITP, and IVIG was administered, which resulted in an increased platelet count of 139 x 10^9/L and Hb > 10.0 g/dl. A month later the platelets dropped 12 x 10^9/L, although her Hb remained at 12.1 g/dl, with the presence of rare schistocytes on blood smears. A second course of IVIG was delivered, and her platelet count rose to 301 x 10^9/L. After 9 months, the patient developed a petechial rash. Her CBC was indicated thrombocytopenia (19 x 10^9/L), low Hb (4.9g/dL), reticulocytosis (4.1%), and the presence of 3-5 schistocytes per high-power view. The patient’s LDH was 1183 U/L, urea was 4.5 mmol/L, and creatinine was 45 μmol/L. The direct Coombs test was negative. ADAMTS13 was 0%, and anti-ADAMTS13 inhibitory antibodies were > 4.5 Bethesda units. The patient was diagnosed with immune TTP, and high dose IV methylprednisolone 500 mg/d was administered for 3 days, and FFP infusions of 20 ml/kg day were started.

The patient was referred to our centre. Upon admission, the patient was fatigued, with extensive cutaneous haemorrhages. The patient’s CBC showed low Hb (9.6 g/dL), reticulocytes (20.5%), schistocytes (2.8%), and low platelets (8 x 10^9/L). The total bilirubin level was 47.2 μmol/L, LDH was 1551 U/L, urea was 6.6 mmol/L, creatinine was 43 μmol/L, and proteinuria was indicated at 2.0 g/L. ADAMTS13 activity was 0%. The patient underwent PEX daily, rituximab 375 mg/m^2 was administered twice weekly for 7 doses. After 8 PEX procedures, the patient’s platelets recovered to > 150 x 10^9/L, and the frequency of the procedure dropped to every other day. Ten days after the last PEX procedure, the patient’s platelets dropped to 90 x 10^9/L, and her Hb decreased to 10.0 g/dL. ADAMTS13 activity was 10%. PEX was resumed, and the patient received bortezomib, 1.3 mg/m^2 on days 1, 4, 8, and 11. After 2 courses of PEX, the patient’s platelets increased to 249 x 10^9/L, and ADAMTS13 activity increased to 24%. After two months, her platelets had stabilised at > 300 x 10^9/L, ADAMTS13 activity had reached 60%, and inhibitors were no longer detectable. Twelve months after the last bortezomib dose, the patient remains free of disease, with normal ADAMTS13 activity and no inhibitors (figure 2).
Discussion

Treatment of immune-mediated TTP with PEX aims to eliminate ADAMT13 inhibitors from circulation, and to provide ADAMTS13 in amounts sufficient for cleaving ULVWF, while immunosuppression aims to stop further production of inhibitors by B-lymphocytes and plasma cells. It has been shown that treatment with rituximab in addition to PEX and glucocorticoids increases the probability of achieving remission, reduces the probability of relapse, and shortens the interval to remission. Relapses of TTP are common and are due to the continued production of anti-ADAMTS13 antibodies. Hematologic remission are not equivalent to immunologic remission, as levels of ADAMTS13 activity exceeding 5-10% are sufficient in a steady state to maintain the appropriate cleavage of ULVWF. Thus, the ultimate goal of TTP treatment is to restore ADAMTS13 activity to normal by eliminating anti-ADAMTS13 antibodies. Historically, many immunosuppressive interventions, such as vincristine, cyclophosphamide, or splenectomy, have been used in patients with TTP with varying degrees of success.

The proteasome inhibitor bortezomib, have proven to be an effective therapy of multiple myeloma, resulting in the long-term suppression of malignant plasma cells. Bortezomib has been used successfully in the treatment of a variety of autoimmune conditions, including TTP, but only one publication has reported its use in adolescents. In line with published cases, our two patients with short-lived responses to PEX, rituximab, and high-dose glucocorticoids subsequently went into long-term remission with the elimination of inhibitors and normalisation of ADAMTS13 activity after just one course of bortezomib. No side effects of bortezomib were seen, indicating that this treatment is safe and thus suitable for use in children. Monitoring the level of ADAMTS13 activity and inhibitor activity after the achievement of clinical and hematological remission is appropriate for determining the timing of this intervention.

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Author contribution:
AM proposed using bortezomib and wrote the text, UP, IK, DF, DB and NK provided clinical care and data retrieval, EK provided plasma exchange, PA assayed ADAMTS13 activity and anti-ADAMTS13 inhibitors titres, GN participated in discussion and provided valuable advices about paper final version.

References


Figure 1.
Therapies delivered and evolution of laboratory parameters of patient 1. Solid line indicates platelet counts, dash line depicts ADAMTS13 activity. PEX and arrows indicate plasma exchanges, R with white arrowheads indicate rituximab, M followed by circles indicate high-dose methylprednisolone, and B with black arrowheads indicate bortezomib.

Figure 2.
Therapies delivered and evolution of laboratory parameters of patient 1. Solid line indicates platelet counts, dash line depicts ADAMTS13 activity. PEX and arrows indicate plasma exchanges, R with white arrowheads indicate rituximab, M followed by circles indicate high-dose methylprednisolone, and B with black arrowheads indicate bortezomib.