Integrating new treatment options into the management of adult ITP

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Abstract
Treatment of steroid resistant ITP in adults can be challenging in patients who are actively bleeding. The majority of novel therapies that have been developed in the last few years including anti CD 20 monoclonal antibody therapy (Rituximab) and the thrombopoietic growth factors romiplostim and eltrombopag, take time to work. Combinations of active agents may accelerate the response rate. Splenectomy and the use of immunosuppressive agents may still have an important role in the acute management. A case of resistant ITP will be discussed in the context of currently available treatment modalities.

Introduction

Immune Thrombocytopenia is an autoantibody mediated blood disorder characterized by both increased platelet destruction and decreased platelet production. The estimated incidence is 9.5 cases per 100,000 population with an estimated 200,000 cases per year in the United States. Median age at presentation is 56 with 2 peaks: age 20-40 and age over 60. There is no male/female predominance in adults. The disorder is self-limited in 70-80% of children.

Immune thrombocytopenia typically presents with bleeding from mucosal surfaces which may include epistaxis, oral mucosal bleeding, or bleeding from the GI or Genitourinary tracts. Critical site bleeding, including the brain, may occur. Spontaneous bruising and a petechial rash are common presenting complaints. Many patients present with an isolated abnormality found on routine laboratory analysis [1-6].

An International consensus panel redefined the older terminology of ITP (Idiopathic Thrombocytopenic Purpura) into Primary and Secondary Immune Thrombocytopenia. Primary immune thrombocytopenia is a diagnosis of exclusion with isolated thrombocytopenia in the absence of other causes or disorders that may be associated with thrombocytopenia. Secondary immune thrombocytopenia comprises all forms of immune thrombocytopenia except primary ITP. Treatment is targeted to the underlying disease or offending medication/drug and usually has a different natural history compared with primary ITP (Table 1 and Figure 1) [2,7,8].

There are no robust laboratory or clinical parameters available to establish the diagnosis with accuracy. The platelet threshold for diagnosis was changed from ≤ 150,000 to ≤ 100,000 based on worldwide epidemiologic observations that there was only a 6.9% 10 year probability that a platelet count in the 100-150,000 range would get worse over time. Testing for the presence of anti-platelet antibodies has not been informative to date [9,10].

Case History

The patient is a 40 year old Hispanic woman, originally from Mexico, in excellent health, on no chronic medications, who presented with the sudden onset of gingival bleeding, menorrhagia, and headache. She did not use alcohol or illicit drugs. She had 6 uncomplicated pregnancies, most recent 3 years prior, and she was still breast feeding. There was no recent travel. No infectious exposures. Previous testing for HIV, RPR, and Hepatitis B surface antigen at time of recent pregnancy was negative. There was no family history of a blood disorder.

Exam
Obese woman 103 kg. Afebrile
Diffuse gum bleeding in the oropharynx. No cardiac murmur. No lymphadenopathy. No organomegaly. No focal neurologic deficits. No retinal hemorrhages. Optic discs sharp Skin with a diffuse petechial rash on the trunk and extremities.

Initial Lab
Hemoglobin 12.3, hematocrit 37%, MCV 85; WBC 5,800 66 neutrophils, 24 lymphocytes, 8 monocytes, 1 eosinophil, platelet count <5,000.
BUN 12, creatinine 0.6
Uncorrected reticulocyte count 1.2%; bilirubin 0.7; LDH 185; haptoglobin 127
Prothrombin time 15.5 seconds (normal up to 15.2); PTT 39 seconds (normal up to 37); fibrinogen 351 mg%; D-dimer <0.27.

Review peripheral blood film
Normochromic, normocytic red cells, no spherocytes, schistocytes, or polychromasia. Less than one platelet per 10 high powered fields.
Platelet count 195,000 7/15/13; 136,000 2/4/14 at time of pregnancy 3 years prior, not repeated since then.

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Clinical Course and Management

This patient presented with both non critical and critical site bleeding. Platelet transfusion indicated in view of critical site bleeding into the brain with early subarachnoid hemorrhage with data showing up to a 40% response [11,12,18].

Corticosteroids remain the mainstay of initial treatment for newly diagnosed ITP: prednisone 1-2 mg/kg/day for 28 days then slowly tapered. An increasing body of data supports the use of pulse, oral, high dose dexamethasone as an alternative [12-14]. In the only prospective, randomized, clinical trial of standard dose prednisone compared with high dose dexamethasone, Yu Wei, et al. demonstrated that dexamethasone 40 mg daily for 4 days resulted in a higher initial response, an approximate doubling of the complete response rate, and a median 50% shorter time to response with less toxicity compared with standard prednisone (Table 2) [15].

In the actively bleeding patient, alternative steroid dosing with methylprednisolone (Solumedrol) in doses of 30 mg/kg up to One gram Intravenous for 7 days has been used in conjunction with intravenous immunoglobulin 1 gram/kg x 2 doses [1,4,17]. Anti-D (Rh) globulin 75 micrograms/kg IV over 20 minutes in non splenectomized patients has also been given in conjunction with steroids. Multiple potential mechanisms of action have been proposed for the efficacy of IVIG and/or anti-D in ITP but the major mechanism is felt to be displacement of antibody coated platelets from Fc receptors in the spleen [17,25]. A bleeding score has been developed to assist in clinical decision making in view of the fact that there is a poor correlation between the absolute platelet count and the propensity to bleed in an individual patient [24].

The patient was given pulse high dose dexamethasone and IVIG. Aygestin, oral estrogen, was initiated to suppress menses. AMICAR was initially given by oral rinse to decrease oral mucosal bleeding and by intravenous to limit intracranial bleeding.

By hospital day 6, there was no platelet response and continued oropharyngeal and vaginal bleeding consistent with steroid refractory, persistent, ITP. Options considered were anti CD-20 monoclonal antibody therapy with Rituximab, thrombopoietin receptor agonists, or splenectomy.

Rituximab targets B lymphocytes resulting in Fc mediated cell lysis with the ability to deplete autoantibody producing B cells active in ITP. There is an overall 40-60% response rate. Responses are only durable in 20-40% of patients. A major limitation in the treatment of acute ITP is the slow median time to response of 5.5 weeks. [25-30]. Rituxan plus high dose dexamethasone has been studied as a component of initial therapy but it is unclear whether it’s use impacts on the rapidity of response. In a randomized, open label, prospective, phase 3 study in 9 centers in Denmark, 137 newly diagnosed patients with primary ITP with platelet counts less than 25,000 or less than 50,000 with active bleeding, were randomized to high dose dexamethasone or dexamethasone plus Rituximab. The primary endpoint was a sustained response. Bleeding more effectively controlled in the HD-DXM arm [15].

Table 1. Differential Diagnosis of ITP. Abbreviations: ITP, idiopathic thrombocytopenic purpura; HIV, human immunodeficiency virus; ESV, Epstein-Barr virus; SLE, systemic lupus erythematosus

<table>
<thead>
<tr>
<th>Immunological</th>
<th>Primary</th>
<th>Secondary</th>
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<tbody>
<tr>
<td>Drug-induced (eg. quinine)</td>
<td>Post-transfusion purpura</td>
<td>HIV</td>
</tr>
<tr>
<td>Post-transfusion purpura</td>
<td>Hepatitis C</td>
<td>Infectious mononucleosis (EBV virus)</td>
</tr>
<tr>
<td>HIV</td>
<td>SLE</td>
<td>Crohn’s disease</td>
</tr>
<tr>
<td>SLE</td>
<td>Antiphospholipid antibody syndrome</td>
<td>Chronic lymphocytic leukemia</td>
</tr>
<tr>
<td>Antiphospholipid antibody syndrome</td>
<td>Lymphoma</td>
<td>IgA deficiency</td>
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<tr>
<td>Chronic lymphocytic leukemia</td>
<td>Common variable immune deficiency</td>
<td>Sarcoïdosis</td>
</tr>
<tr>
<td>Lymphoma</td>
<td>Non-immune</td>
<td></td>
</tr>
<tr>
<td>IgA deficiency</td>
<td>Hyperesplenism</td>
<td>Myelodysplasia</td>
</tr>
<tr>
<td>Common variable immune deficiency</td>
<td>Acute leukemia</td>
<td></td>
</tr>
<tr>
<td>Sarcoïdosis</td>
<td>Drug-induced marrow suppression (valproic acid, alcohol)</td>
<td>Hereditary thrombocytopenia (MYH-9 mutations)</td>
</tr>
</tbody>
</table>

Microangiopathic hemolytic anemia

Table 2. Initial CR was a positive indicator of sustained response. 116 patients (60%) had baseline platelet count ≤ 10,000. HD-DXM arm with consistently higher counts after initial response. Bleeding more effectively controlled in the HD-DXM arm [15]

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Prednisone (n=98)</th>
<th>High Dose Dex (n=97)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Response</td>
<td>67%</td>
<td>82%</td>
</tr>
<tr>
<td>Overall response</td>
<td>74%</td>
<td>85%</td>
</tr>
<tr>
<td>Complete Response</td>
<td>26.8%</td>
<td>50.5%</td>
</tr>
<tr>
<td>Time to response(days)</td>
<td>6(2-24)</td>
<td>3(1-9) p&lt;0.001</td>
</tr>
<tr>
<td>Sustained response</td>
<td>41.2%</td>
<td>40%</td>
</tr>
</tbody>
</table>

Figure 1. Estimated fraction of the various forms of secondary ITP based on clinical experience of the authors. ALPS indicates autoimmune lymphoproliferative syndrome; posttx, post–bone marrow or solid organ transplant; SLE, systemic lupus erythematosus; APS, antiphospholipid syndrome; CLL, chronic lymphocytic leukemia; CVID, common variable immune deficiency. In the absence of a systematic analysis of the incidence of secondary ITP, the data shown represent our combined assessment based on experience and reading of the literature. Professional illustration by Paulette Dennis [8]

Ancillary Lab Studies

HIV negative, Hepatitis C antibody and quantitative RNA not detectable; Hepatitis B surface antigen negative, surface antibody positive, e-antibody positive, e-antigen negative in an unvaccinated patient reflecting previous resolved hepatitis B infection.

TSH 3.9; urine pregnancy test negative

Imaging

Chest X-Ray: no infiltrate or effusion; no adenopathy

CT brain: bilateral curvilinear areas of hyper-attenuation posterior parietal lobes consistent with low grade subarachnoid hemorrhage.
platelet count of $\geq 50,000$ at 6 months which was achieved in 58% of patients in the combined therapy group versus 37% for dexamethasone alone. Time to response data not reported [31].

A meta-analysis by Chugh et al reviewed 5 eligible trials including 463 non splenectomized patients with ITP who received Rituxan as part of their treatment. In two of the trials, the patients had received no prior treatment. Time to response data not reported. Although complete response rates were higher in the Rituximab treated patients, severe bleeding events did not differ between the two groups [32].

In a small phase Ib study, Choi et al reported that the combination of dexamethasone, low dose Rituximab, and Cyclosporin A given for a single 28 day course resulted in a high response rate when used as first line therapy. 7 of the 20 patients were previously untreated. Overall response was 60% at 6 months. 76% of responding patients were relapse free at 24 months. Median time to response was 7.4 days. However, only 3 of 7 patients with newly diagnosed or persistent ITP responded [33].

Thrombopoietin receptor agonists are assuming a major role in the treatment of relapsed refractory ITP. They are the only agents that have been tested in large, randomized, prospective, placebo controlled trials in ITP. Thrombopoietin is the physiologic stimulator of megakaryocyte growth and differentiation acting through C-mpl, the thrombopoietin receptor (Figure 2). Suboptimal platelet production as well as peripheral destruction of platelets combine to cause the thrombocytopenia of ITP. Mechanisms of suboptimal platelet production include: antibodies directed against immature megakaryocytes resulting in decreased platelet production, increased clearance of endogenous thrombopoietin on antibody coated platelets, and inappropriate feedback rise in TPO levels in the face of platelet destruction [16,34-39].

Current American Society of Hematology Guidelines published in 2011 recommend TPO receptor agonists for patients at risk for bleeding after splenectomy (Grade 1B), or to consider use in patients at high risk of bleeding who have failed one line of therapy and who have not had a splenectomy (Grade 2C) [10].

In practice, given the long term efficacy and safety profile of these agents, they are being used more and more before consideration of splenectomy. A drawback to use in the acute bleeding patient is time to response which is on the average of 1-2 weeks. In the clinical trials that led to the approval of Romiplostim, only 25% of patients responded to the FDA approved starting dose of 1 microgram/kg at 1 week with 50% showing a response after 2 weeks following first dose increase to 2 micrograms/kg [34-42,6,16,19].

Is there a potential to combine TPO receptor agonists with other modalities to accelerate the response rate? Preliminary data using a combination of high dose dexamethasone and Eltrombopag as initial therapy...
therapy are promising. A 100% overall response rate with 83% complete responders by 34 days with 75% ongoing response at 6 months was reported by Gomez-Almaguer [43]. Combinations of TPO receptor agonists with immunosuppressive agents are also under investigation. In a multi-center, randomized, open label study by Zhou, et al. 115 patients with primary ITP and platelet counts <30,000 who had failed either steroids, splenectomy, or cyclosporine and were off treatment for at least 4 weeks, were randomized to Rituximab fixed dose of 100 mg weekly x 4 or the combination of Rituximab and thrombopoietin 300 micrograms subcutaneously daily x 14 days. Median time to response with the combination was 7 days (4-28), compared with 28 days (4-90) on the Rituxan only arm. Overall response rate 79% vs 71% (p=0.36), CR rate 45 vs 24% (p=0.026) [44] (Table 3).

On hospital day 6, the patient was treated with a combination of Rituximab 375 mg/m² weekly x 4 and Romiplostim 1 mcg/kg SQ with weekly dose escalation to 2 mcg/kg at day 8 then 4 mcg/kg at day15. By day 18, she had persistent bleeding, increased headache, new onset tinnitus, and blurred vision and no platelet response.

Splenectomy was performed on day 19. There was no increase in platelet count with intra and post op platelet transfusion.

Splenectomy remains the gold standard for treatment of refractory ITP with an approximate 80% overall response rate and a 50% durable response past 5 years. Improvement in surgical techniques with the advent of laparoscopic splenectomy has significantly reduced the morbidity and length of hospital stay. Average time to platelet response is 5-7 days [5,6,10,19,38,44-50] [Table 4].

Oral and vaginal bleeding subsided but she complained of increasing tinnitus and headache. Fundoscopic exam now showed bilateral retinal hemorrhages. CT of the brain done on day 23 showed new, subtle, multifocal areas of hemorrhage. K-centra 4 factor clotting was normal. By day 22, there was no improvement in her clinical status. She had 24 interval study with subtle areas of increased hemorrhage. A platelet transfusion was given followed immediately by a dose of IV recombinant VIIa. Parenteral AMICAR dose increased. Another course of high dose dexamethasone was initiated [51].

Immunosuppressive drugs still have a role, albeit limited, in acute management of refractory ITP. There are no large, randomized studies of any agent. Commonly used drugs include 6-mercaptopurine, the calcineurin inhibitors Cyclosporin-A and Tacrolimus, alkylating agents such as Cyclophosphamide, vinca alkaloids, vincristine and vinblastine, other immunosuppressive agents such as Azathioprine, and drugs which work by unclear mechanisms such as Danazol and Dapsone. Responses are seen on the average of 50% of the time. Time to response is, again, an issue with shortest time to response reported for the vincas (5-7 days), followed by cyclophosphamide (7-112 days) (Table 5) [6,38,41,52,53].

On Day 23 Vincristine 2 mg IV was administered.

Bone marrow biopsy showed no additional pathology. Megakaryocytes were present.

On day 25, platelet count began to rise to 7,000: 20 days post initiation of Rituxan/Romiplostim and 7 days post splenectomy, and continued to rise to over one million over the next 7 days with subsequent stabilization at 400,000.

**Discussion**

Treatment of the adult patient with acute Primary Immune Thrombocytopenia who is actively bleeding remains a challenge. Combined modality treatment incorporating novel agents is a promising area of study. The ability to achieve a rapid response is crucial. Current literature demonstrates that more rapid and complete responses can be obtained with the use of high dose dexamethasone as initial therapy combined with Intravenous immunoglobulin. Preliminary study also suggests that the combined use of anti CD-20 antibody therapy and thrombopoietin or Thrombopoietin receptor agonists can also

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**Table 3. Responses and outcomes in the RTX + TPO and RTX groups [44]**

<table>
<thead>
<tr>
<th>Response</th>
<th>RTX + rTPO</th>
<th>RTX</th>
<th>P</th>
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<tbody>
<tr>
<td>CR, % (n)</td>
<td>45.4 (35)</td>
<td>23.7 (9)</td>
<td>0.026*</td>
</tr>
<tr>
<td>PR, % (n)</td>
<td>33.8 (26)</td>
<td>47.4 (18)</td>
<td>0.22</td>
</tr>
<tr>
<td>OR, % (n)</td>
<td>79.2 (61)</td>
<td>71.1 (27)</td>
<td>0.36</td>
</tr>
<tr>
<td>NR, % (n)</td>
<td>20.8 (16)</td>
<td>28.9 (11)</td>
<td>0.36</td>
</tr>
</tbody>
</table>

For patients with an initial response

<table>
<thead>
<tr>
<th>Duration of response</th>
<th>RTX + rTPO</th>
<th>RTX</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>TTR (days), median (range)</td>
<td>7 (4-28)</td>
<td>28 (4-90)</td>
<td>0.004†</td>
</tr>
<tr>
<td>Duration of response (≥ 6 mo), % (n)</td>
<td>67.2 (41)</td>
<td>55.6 (15)</td>
<td>0.34</td>
</tr>
<tr>
<td>Duration of response (≥ 12 mo), % (n)</td>
<td>44.3 (27)</td>
<td>29.6 (8)</td>
<td>0.24</td>
</tr>
<tr>
<td>Duration of response (≥ 24 mo), % (n)</td>
<td>24.6 (15)</td>
<td>18.5 (5)</td>
<td>0.59</td>
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<tbody>
<tr>
<td>* P &lt; 0.05</td>
<td>† P &lt; 0.01</td>
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</tbody>
</table>

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**Table 4. Rx Persistent/Chronic ITP [58]**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Time to Respond</th>
<th>Rate of Response</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Splenectomy</td>
<td>1-24 days</td>
<td>80%</td>
<td>60% sustained over 5-10 yrs</td>
</tr>
<tr>
<td>Danazol</td>
<td>3-6 mos</td>
<td>67% CR/PR</td>
<td>46% at 10 yrs</td>
</tr>
<tr>
<td>IVG</td>
<td>24-48 hrs</td>
<td>80%</td>
<td>2-4 wks</td>
</tr>
<tr>
<td>Anti-D IgG</td>
<td>24-48 hrs</td>
<td>93% initial</td>
<td>43% off Rx &gt;6 mos</td>
</tr>
<tr>
<td>Rituxan</td>
<td>1-8 wks</td>
<td>60% OR</td>
<td>15-20% &gt;3-5 yrs</td>
</tr>
<tr>
<td>TPO Receptor Agonists</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eltrombopag</td>
<td>7-15 days</td>
<td>&gt;80%</td>
<td>Up to 5-6 yrs</td>
</tr>
<tr>
<td>Romiplostim</td>
<td>7-28 days</td>
<td>79-88%</td>
<td>Up to 6 years</td>
</tr>
</tbody>
</table>
accelerate the response rate to treatment. More traditional approaches including splenectomy and immunosuppressive agents still have an important role in individual patients. It is unclear whether the early addition of Rituxan plus romiplostim impacted in a positive way on the time to response to splenectomy cited in the literature is 5 to 7 days [28,29,36,49,50].

A number of new areas of investigation are actively being considered or are under study [6,22,38,40-42,54,55].

Since Romiplostim and Eltrombopag act at different sites on the thrombopoietin receptor, combined use could potentially potentiate and accelerate the thrombopoietic effects.

Immunosuppressive drugs that selectively kill plasma cells such as bortezomib, have shown activity in TTP and might be studied in other immune disorders such as ITP.

Novel immunotherapeutics such as fostanitinib targeting Syk (spleen tyrosine kinase) look promising. Syk is a cytoplasmic tyrosine kinase which is activated when Fc receptors bind to their ligands and promotes phagocytosis of FcγR bound platelets by splenic macrophages. Platelet destruction in ITP is mediated by Syk dependent phagocytosis. Fostanitinib has recently received approved by the FDA in April, 2018 for refractory ITP. C reactive protein also appears to play an important role in individual patients. It is unclear whether the early addition of combined modality therapy can result in rapid responses compared with traditional treatments. This patient failed to respond to initial treatment with high dose dexamethasone and intravenous immunoglobulin and continued to bleed. Early addition of combined weekly anti-CD 20 antibody therapy at a standard dose of 375 mg/ meter squared and a Thrombopoietin receptor agonist started at 1 microgram/kg with weekly dose escalation, did not result in a response by day 13 after initiation and splenectomy was required. There was a delayed response to splenectomy prompting the addition of a single dose of vincristine on day 23. Response was seen on day 25, 20 days after initiation of Rituxan/Romiplostim and 7 days post splenectomy.

Although combined modality therapy, particularly the use of high dose dexamethasone and a TPO agonist with or without Rituximab appears promising, more prospective data is needed to justify use in the patient who presents with major bleeding and early splenectomy remains the standard of care.

References


44. Zhou H, Xu M1, Qin F, Zhang HY, Yuan CL, et al. (2015) A multicenter randomized open-label study of rituximab plus rhTPO vs rituximab in corticosteroid-resistant or relapsed ITP. Blood 125: 1541-1547. [Crossref]


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