Eltrombopag for the treatment of aplastic anemia: current perspectives

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Abstract: Aplastic anemia (AA) is a potential life-threatening hematopoietic stem cell (HSC) disorder resulting in cytopenia. The mainstays of treatment for AA are definitive therapy to restore HSCs and supportive measures to ameliorate cytopenia-related complications. The standard definitive therapy is HSC transplantation for young and medically fit patients with suitable donors and immunosuppressive therapy (IST) with antithymocyte globulin and cyclosporine for the remaining patients. A significant proportion of patients are refractory to IST or relapse after IST. Various strategies have been explored in these patients, including second course of antithymocyte globulin, high-dose cyclophosphamide, and alemtuzumab. Eltrombopag, a thrombopoietin mimetic, has recently emerged as an encouraging and promising agent for patients with refractory AA. It has demonstrated efficacy in restoring trilineage hematopoiesis, and this positive effect continues after discontinuation of the drug. There are ongoing clinical trials exploring the role of eltrombopag as a first-line therapy in moderate to severe AA and a combination of eltrombopag with IST in severe AA.

Keywords: eltrombopag, aplastic anemia, thrombopoietin, c-Mpl receptors

Introduction
Aplastic anemia (AA) is a potential life-threatening hematopoietic stem cell (HSC) disorder with an estimated incidence of 2–3 per million per year.1,2 The incidence is triphasic, with the first peak occurring in patients at 2–5 years of age, the second peak at 20–25 years, and the third peak at >60 years of age. It is characterized by pancytopenia with a hypocellular marrow in the absence of abnormal infiltration or marrow fibrosis.3–5 Its diagnosis requires exclusion of clonal disorders, including myelodysplastic syndrome (MDS), hypocellular myeloid leukemia, and paroxysmal nocturnal hemoglobinuria. Its severity depends on the degree of peripheral cytopenias and the marrow cellularity (Table 1). Severe AA (SAA) is defined as bone marrow cellularity of <25% with two out of three of the following: neutrophils <0.5×10^9/L, platelet count <20×10^9/L, and reticulocyte count <20×10^9/L. The presenting features range from an incidental finding of cytopenias on a routine blood count to symptomatic cytopenias, including pallor, bleeding, and infection. AA is associated with substantial morbidity, leading to reduced quality of life and premature death. Prolonged neutropenia increases susceptibility to potentially fatal infections, while thrombocytopenia might result in catastrophic bleeding in major organs. Chronic iron overload secondary to frequent red cell transfusion inevitably leads to multisystemic dysfunctions that encompass cardiomyopathy, endocrinopathies, and liver siderosis.

The etiology of AA can be broadly classified into inherited AA and acquired AA. Inherited AA is rare, and the diagnostic clues to this are skeletal abnormalities,
skin pigmentary anomalies, and marrow dysplasia. Absence of these features does not rule out inherited disorders, and genetic testing is recommended to exclude marrow failure syndrome, as it has critical impact on clinical management and donor selection for HSC transplantation (HSCT). The four important causes for inherited AA are Fanconi anemia (FA), dyskeratosis congenita, Shwachman–Diamond syndrome, and congenital amegakaryocytic thrombocytopenia (CAMT). FA is the most common of inherited AA. Although most patients with FA present in childhood, the diagnosis might be delayed until adulthood, as characteristic congenital malformations are absent in 30%–40% of patients. Hence, FA should be considered as a part of workup for patients with AA. HSCT is the only current curative treatment for inherited AA, and these patients require specific transplant considerations to prevent unexpected toxicities.

Acquired AA accounts for ~80%–90% of marrow failure. It is distinguished from constitutional marrow failure by the absence of phenotypic features and the absence of mutation in genes known to be responsible for inherited AA. The causes of acquired stem cell failure include direct stem cell destruction by drugs, chemicals, ionizing radiation, and viruses, but no identifiable cause is found in the majority of patients. The pathology underpins the marrow failure in idiopathic AA is immune dysfunction, resulting in paucity of HSCs. The most convincing evidence supporting the immune-mediated destruction in idiopathic AA is its response to immunosuppressive therapy (IST) and rebound cytopenia with discontinuation of IST. The proposed hypothesis is that there is an aberrant T-cell activation stimulated by a viral infection, a drug, or a specific environmental precipitant in a genetically susceptible host. Clonal expansion of CD8+ T-cells and Th1-cells, reduced and dysfunctional regulatory T-cells

### Table 1 Overview of aplastic anemia

| Definition | Pancytopenia with hypocellular marrow in the absence of abnormal infiltration or marrow fibrosis |
| Incidence | 2–3 per million per year |
| Age | Triphasic distribution: 2–5 years, 20–25 years, and >60 years |
| Diagnostic criteria | Any two of the following features: |
| | 1. Hemoglobin <100 g/L |
| | 2. Platelets <50×10^9/L |
| | 3. Neutrophils <1.5×10^9/L |
| Severity | Marrow cellularity <25% with any two of the following features: |
| | 1. Neutrophils <0.5×10^9/L |
| | 2. Platelets <20×10^9/L |
| | 3. Reticulocyte count <20×10^9/L |
| Causes | As above plus neutrophils <0.2×10^9/L |
| Inherited AA | Not fulfilling the criteria for severe of very severe AA |
| 1. FA | Acquired AA |
| 2. DKC | 1. Idiopathic (70%–80%) |
| 3. SDS | 2. Posthepatitis |
| 4. CAMT | 3. Postviral infection |
| Causes | 4. Drugs |

### Diagnostic evaluation of AA

1. Full blood count |
2. Reticulocyte count |
3. Blood smear |
4. Bone marrow aspiration and trephine biopsy |
5. HbF |
6. Peripheral blood chromosomal breakage analysis |
7. Flow cytometry for GPI-anchored proteins to detect PNH clone |
8. Liver function tests |
9. Viral studies: hepatitis A/B/C, EBV, CMV, HIV, and parvovirus B19 |
10. Vitamin B12 and folate |
11. Autoimmune screening |
12. Imaging: echocardiogram and abdominal ultrasound scan |
13. Next-generation sequencing gene panel for marrow failure syndrome |

**Abbreviations:** AA, aplastic anemia; CAMT, congenital amegakaryocytic thrombocytopenia; DKC, dyskeratosis congenita; FA, Fanconi anemia; PNH, paroxysmal nocturnal hemoglobinuria; SDS, Shwachman–Diamond syndrome; HbF, hemoglobin F; GPi, glycosylphosphatidylinositol; EBV, Epstein-Barr virus; CMV, cytomegalovirus.
(Tregs), and increased Th2- and Th17-cells are the hallmarks of immune dysregulation. The hemostasis of hematopoiesis is interrupted by marrow-suppressing cytokines, including interferon gamma and tumor necrosis factor. The apoptotic death of HSCs in the bone marrow is mediated by increased expression of the Fas receptor and antigen, as well as possibly by cytotoxic granules.

Current recommendation of management of patients with acquired SAA is illustrated in Figure 1. In general, patients with SAA need definite therapy to target the marrow failure and support care for cytopenia-associated clinical symptoms and complications. The components of the support care of AA encompass transfusion supports, antimicrobial prophylaxis, treatment for neutropenic fevers, and hematologic growth factor therapies. The definite therapy for SAA includes HSCT and IST. Both are effective and aim to abrogate the immune-mediated depletion of HSC. By stopping the immune destruction of HSC, IST allows repopulation of the marrow from the surviving, depleted HSC pool. In HSCT, administration of donor-derived HSC allows more rapid restoration of blood counts, and there is also a reduced incidence of later clonal hematologic disorders than following IST. Matched sibling donor HSCT has always been the therapy of choice for young patients with SAA. The event-free survival with human leukocyte antigen-matched sibling donor transplant is 75%–90%, and the risk of graft failure is 4%–14% using conditioning regimen with cyclophosphamide and antithymocyte globulin. Transplant-related morbidity, such as graft-versus-host disease and infection, has reduced significantly with better transplant care. There are few later effects since the conditioning therapy is light and growth and puberty are preserved. In addition, with better unrelated donor availability, the outcome of well-matched unrelated donor HSCT in all indications is approaching matched sibling donor. Some institutions and national groups are using matched unrelated donor HSCT as the preferred initial therapy rather than IST for young patients.

With regard to IST, the standard regimen is a combination of antithymocyte globulin and cyclosporine. The response to IST is delayed and usually seen after 3–4 months. Of those patients treated with IST, 20%–30% do not respond while 10%–30% of responders relapse. The mechanism

**Figure 1** Current recommendation for management of severe aplastic anemia.

**Abbreviations:** AA, aplastic anemia; ATG, antithymocyte globulin; HLA, human leukocyte antigen; HSCT, hematopoietic stem cell transplantation; IST, immunosuppressive therapy; SAA, severe AA.
supports the relapse, and refractory diseases might be due to an uncontrolled immune dysfunction or deficiency in residual HSC. Various strategies, including eltrombopag, alentuzumab, and high-dose cyclophosphamide, have been proposed for these patients. Patients treated with IST are at risk of developing clonal evolution, including MDS/acute myeloid leukemia (8%), hemolytic paroxysmal nocturnal hemoglobinuria (10%), and solid tumor (11%).

The article describes the principle of eltrombopag use in acquired AA and summarizes the current evidence for its use in AA.

**The role of thrombopoietin in hematopoeisis**

Thrombopoietin (TPO) is the most important hematopoietic growth factor that regulates the circulating platelet mass. Although the concept of this platelet regulator was first described in 1950, TPO protein was purified only in 1994. TPO is also known as megapoietin, megakaryocyte growth and developmental factor, and c-Mpl ligand. The TPO gene is located on the long arm of chromosome 6 at the position 3q27–28. It has seven exons extending along ~7,000 bp, and the first two exons are noncoding. TPO is mainly produced by liver, with small amounts being made by the kidney and bone marrow. It consists of 353 amino acids with 21-amino-acid secretory leader sequence. The mature TPO protein is a member of four-helix-bundle cytokine superfamily and consists of two domains. The amino-terminal 154 residue domain is homologous to erythropoietin and binds to the c-Mpl receptor. The main two functions of carboxyl-terminal domain of TPO are serving as an intramolecular chaperone to aid the proper folding of the polypeptide into the mature hormone and prolonging the circulatory half-life of TPO by modifying with multiple sites of both N- and O-linked carbohydrates.

TPO is a potent endogenous cytokine that acts through the TPO receptors, known as c-Mpl receptors, which present primarily in platelets and megakaryocytes and in a small percentage of hematopoietic progenitor cells (HPC). The c-Mpl gene, which is located on human chromosome 1p34, was cloned in 1992. c-Mpl protein exists as an inactive dimer, each monomer containing two cytokine receptor homology (CHR) domains. Binding of TPO to distal cytokinecytokine receptor homology region of c-Mpl receptors stimulates multiple signal transduction pathways, including JAK/STAT and mitogen-activated protein kinase pathways. Activation of these pathways promotes megakaryocyte proliferation and maturation, as well as platelet release into circulation.

TPO clearance depends on its binding to c-Mpl receptors. Increasing free TPO in thrombocytopenia state leads to stimulation of platelet production, whereas its level is low in thrombocytosis. As c-Mpl receptors are present in HPC, TPO has been shown to play an important role in HSC survival, self-renewal, and expansion. Mutations in the c-Mpl gene have been reported in association with familial AA, and patients with c-Mpl-associated CAMT are at high risk of developing AA. Deficiency of c-Mpl receptors in mouse model demonstrated reduced numbers of HPC. This evidence supports the role of TPO beyond its primary function as a platelet regulator.

With the discovery of TPO, a number of TPO receptor agonists or TPO mimetics have been developed (Table 2). The two, first generation of TPO mimetics, recombinant human TPO (rHuTPO) and PEGylated rHuTPO (PEG-rHuMGDF), were introduced in 1997. rHuTPO shares the same amino acid structure as endogenous TPO but a slightly lower molecular weight. rHuTPO was administered intravenously with a half-life of 30–40 hours. PEG-rHuMGDF was composed of the amino-terminal 163 amino acids of endogenous TPO coupled with polyethylene glycol moiety. It had a half-life of 25–35 hours after an intravenous injection. Although both have demonstrated positive platelet increment, clinical experience with the first-generation TPOs was disappointing because of the development of endogenous antibodies against endogenous TPO, resulting in secondary thrombocytopenia. Developments of PEG-rHuMGDF and rHuTPO were stopped in 1998 and 2002, respectively.

The experience from the first generation of TPO mimetics has formed an important learning platform for the new generation of thrombopoietic growth factors. The second-generation molecules, including small molecular peptides and nonpeptide agents, were first introduced in 1997. Romiplostim is an injectable peptide TPO mimic that activates the TPO receptors just like native TPO. It is composed of 14-amino-acid peptides attached to two disulfide-bonded human immunoglobulin G1 kappa heavy chain constant regions by glycine bridges. As the peptide has no sequence homologous with endogenous TPO, production of antibody against romiplostim would not cross-react with endogenous TPO. It has been used in patients with immune thrombocytopenia (ITP), MDS, chemotherapy-induced thrombocytopenia, and hepatitis C-related thrombocytopenia. Nonpeptide TPO receptor agonists have been derived from small molecule screening techniques that uncovered a number of chemical compounds that interact with c-Mpl receptors. Eltrombopag,
Table 2 Key milestones in eltrombopag

<table>
<thead>
<tr>
<th>Year</th>
<th>Event</th>
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<tbody>
<tr>
<td>1950</td>
<td>First description of platelet growth factor</td>
</tr>
<tr>
<td>1994</td>
<td>TPO protein was purified</td>
</tr>
<tr>
<td>1995</td>
<td>First generation of thrombopoietic mimetics: rHuTPO and PEG-rHuMGDF</td>
</tr>
<tr>
<td>1997</td>
<td>Second generation of thrombopoietic mimetics: eltrombopag and romiplostim</td>
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<tr>
<td>2004</td>
<td>Phase I clinical study of eltrombopag</td>
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<tr>
<td>2007</td>
<td>First Phase II clinical trial of eltrombopag in adults with chronic ITP</td>
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<tr>
<td>2007</td>
<td>First Phase II study of eltrombopag for thrombocytopenia in adults with cirrhosis associated with hepatitis C</td>
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<tr>
<td>2011</td>
<td>Phase III study of eltrombopag in adults with chronic ITP (RAISE)</td>
</tr>
<tr>
<td>2012</td>
<td>First Phase II clinical trial of eltrombopag in refractory AA</td>
</tr>
<tr>
<td>2013</td>
<td>EXTEND study on safety and efficacy of long-term eltrombopag in the treatment of adults with chronic ITP</td>
</tr>
<tr>
<td>2015</td>
<td>First clinical trial of eltrombopag for the treatment of children with persistent and chronic immune thrombocytopenia (PETIT)</td>
</tr>
<tr>
<td>2015</td>
<td>Multicenter, randomized, placebo-controlled, double-blind, Phase I/III trial of eltrombopag in patients with advanced myelodysplastic syndrome and acute myeloid leukemia</td>
</tr>
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</table>

Ongoing clinical trials

- NCT01703169: Efficacy and safety of eltrombopag in patients with severe and very severe AA
- NCT01328587: A pilot study of a TPO-R agonist, eltrombopag, in moderate AA patients
- NCT01891994: Extended dosing with eltrombopag in refractory severe AA
- NCT01623167: Ebtrombopag added to standard immunosuppression in treatment-naive severe AA
- NCT02099747: A prospective randomized multicenter study comparing hATG + CsA with or without eltrombopag as front-line therapy for severe AA patients

Abbreviations: AA, aplastic anemia; CsA, cyclosporine A; hATG, horse antithymocyte globulin; ITP, immune thrombocytopenia; PeG-rHuMGDF, PeGylated rHuTPO; rHuTPO, recombinant human TPO; TPO, thrombopoietin; TPO-R, TPO-receptor.

which has been derived from hydrazinonaphthalene, has a number of advantages compared to peptide TPO mimetics, including oral preparation and a different mechanism of action that confers additive effect. Eltrombopag is the only orally available nonpeptide TPO receptor agonist approved by the US Food and Drug Administration for the treatment of adults with ITP, patients with hepatitis C-related thrombocytopenia who are being treated with interferons, and patients with AA who are refractory to IST.47

Pharmacology of eltrombopag

Eltrombopag is a member of the biarylhydrazone class of compound with a chemical structure of C_{28}H_{27}N_{3}O_{4}. In clinical studies, it has been used as eltrombopag olamine, which is the bis-monoethanolamine salt form of eltrombopag.28 It is rapidly absorbed following oral administration and achieves peak plasma concentrations at a median time of 2.5 hours (Table 3). The biological half-life is ~21–32 hours, and it is mainly excreted in feces and urine. It is highly bound (>99%) to plasma proteins, mainly albumin, and extensively metabolized by monoxygenation, glucuronidation, hydrazine cleavage, and secondary oxidation and conjugation. In vitro, it is an inhibitor of CYP2C8 and CYP2C9, several glucuronosyltransferases isoenzymes, breast cancer resistance protein, and organic anion transporting polypeptide (OATP0 1B1).28,44

In vitro studies have demonstrated that the activity of eltrombopag is dependent on the expression of TPO receptors.49 It binds to the transmembrane domain of the TPO receptor, leading to signal transduction through various pathways, including Janus kinase/signal transducer and activator of transcription and mitogen-activated protein kinase. Activation of TPO receptor results in proliferation and differentiation of megakaryocytes and increased platelet production. As eltrombopag binds at transmembrane domain, it does not compete with endogenous TPO binding at the extracellular TPO receptor domain (Figure 2).49 Hence, eltrombopag provides an additive effect with endogenous TPO in promoting megakaryocyte maturation and platelet production.

Studies using eltrombopag in chronic ITP have demonstrated that prolonged administration of eltrombopag is safe and well tolerated.50–53 Similar to other TPO mimetics, the potential toxicities of eltrombopag include thrombocytosis, thrombosis, reversible bone marrow fibrosis, rebound thrombocytopenia, cataract formation, and reversible hepatic dysfunction. The proposed mechanisms for thrombosis of TPO mimetics are rapid elevation of platelet count and increased rate of microparticle formation in patients with ITP.54 Eltrombopag might be associated with a risk of portal vein thrombosis in patients with advanced liver disease.
Table 3 Drug property of eltrombopag

<table>
<thead>
<tr>
<th>Generic drug name</th>
<th>Eltrombopag olamine</th>
</tr>
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<tbody>
<tr>
<td>Route of administration</td>
<td>Oral</td>
</tr>
<tr>
<td>Preparation</td>
<td>12.5 mg, 25 mg, 50 mg, 75 mg, and 100 mg</td>
</tr>
<tr>
<td>Chemical structure</td>
<td>C_{25}H_{22}N_{4}O_{4}</td>
</tr>
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Mechanism of action
TPO nonpeptide agonist that binds to the transmembrane and iuxtamembrane domain of TPO receptor. It activates intracellular signal transduction pathways, JAK/STAT, and mitogen-activated protein kinase to increase the proliferation and differentiation of HSC

Pharmacokinetic data
- **Bioavailability**: ~52%
- **Protein binding**: >99%
- **Metabolism**: Extensive hepatic metabolism via CYP 1A2, 2C\(^+\) oxidation and UGT 1A1 and 1A3 glucuronidation
- **Biological half-life**: ~21–32 hours
- **Time to peak, plasma**: 2–6 hours
- **Excretion**: Feces (~59%) and urine (~31%)

Clinical use
1. Chronic immune thrombocytopenia
2. Chronic hepatitis C-associated thrombocytopenia
3. Refractory severe aplastic anemia

Potential risks
1. Thrombosis
2. Bone marrow fibrosis
3. Clonal evolution
4. Rebound thrombocytopenia
5. Antibody formation
6. Cataract

Common reported side effects (>10%)
1. Reversible hepatic dysfunction
2. Headache
3. Gastrointestinal symptoms (anorexia, vomiting, diarrhea, and abdominal pain)
4. Pyrexia
5. Fatigue
6. Cough
7. Alopecia
8. Arthralgia and myalgia

Abbreviations: HSC, hematopoietic stem cell; TPO, thrombopoietin.

Of 142 patients with chronic liver disease treated with eltrombopag, seven patients developed portal vein thrombosis compared to three patients in the control group \((n=147)\) (odds ratio 3.04, 95% CI 0.62–14.82).\(^5\) Similar association was not seen in 74 patients with hepatitis C treated with eltrombopag.\(^5\) The use of eltrombopag in ITP did not increase the rate of thrombosis compared to placebo.\(^5\)

With regard to marrow fibrosis, mouse models demonstrated higher fibrosis after TOP gene transfection and reversible fibrosis after discontinuation of TPO.\(^5\) In a recent paper by Brynes et al, of 232 marrow biopsy specimens taken from 117 patients with chronic ITP treated with eltrombopag in the EXTEND study, two patients were found to have moderate to marked reticulin fibrosis. After discontinuing the treatment, the repeat biopsy was normal in one patient and unchanged in another patient. None of the 117 patients developed MDS and acute myeloid leukemia.\(^6\) Similar findings of reversible marrow fibrosis were reported by Kuter et al on 271 patients with chronic ITP treated with romiplostim. Although there is no long-term prospective bone marrow evaluation in patients treated with eltrombopag, the reported studies of eltrombopag in AA and post-HSCT thrombocytopenia did not show evidence of marrow fibrosis in their cohorts.\(^5\)\(^,\)\(^6\)

The experience on the use of eltrombopag in children is limited, and most of the studies are on adults with chronic ITP. PETIT, the first international trial on eltrombopag for the treatment of children with persistent and chronic ITP, identified the dose to be used in Phase III studies.\(^6\) PETIT2 was
done to study the efficacy of eltrombopag in children with chronic ITP aged 6–17 years. In these studies, there were no unexpected safety concerns, and few children discontinued treatment because of adverse effects.51

**Eltrombopag in AA**

The thrombocytopenia in AA is caused by inadequate HSC numbers and function, which lead to impaired megakaryopoiesis and reduced mature platelet production. The consequences of thrombocytopenia in these patients range from silent hemorrhage, such as minor cerebral bleeds, to troublesome mucocutaneous hemorrhage and fatal major organ hemorrhage. The mainstay of therapy for thrombocytopenia in these patients is prophylactic and therapeutic platelet transfusion while waiting for definitive therapy. Antibody-mediated platelet refractoriness is the momentous sequela from multiple donor exposure, although its incidence has reduced from 50% to 12% with the use of leukodepleted blood products.64 The use of hematopoietic growth factor (HGF) in correcting bone marrow failure syndrome has been explored since 1960. The early clinical trials using erythropoietic growth factors, G-CSF, GM-CSF, stem cell factor, interleukin (IL)-1, IL-3, and IL-6 have failed to show beneficial effects. The lack of positive therapeutic effects of these growth factors might be explained by: 1) normal mRNA expression and secretion of these HGFs and cytokines in long-term marrow cultures from AA stromal cells, 2) marked elevation of serum HGF in patients with AA, and 3) activity of these cytokines on more committed myeloid progenitors.65

In addition, significant toxicities were reported with most agents, except G-CSF.65 As TPO receptors, c-Mpl receptors present on more primitive HSCs, which supports the ability of TPO to act on less differentiated progenitor cells and the theoretical concept of treating AA.

As discussed earlier, knockout mice that are deficient in TPO and c-Mpl receptors have reduced numbers of multiple hematopoietic lineage progenitor cells.33,34 Similarly, patients with CAMT are associated with c-Mpl mutation and develop AA at a median age of 3.7 years.66 These findings have reaffirmed the role of TPO and its receptors in multilineage hematopoiesis.

Although the clinical trials of eltrombopag are limited, they have demonstrated the efficacy of the drug in increasing platelet count in patients with chronic ITP and thrombocytopenia associated with hepatitis C infection. The first and only published clinical trial of eltrombopag in AA was performed by Olnes et al in 2012 (Figure 3). In this Phase II study, 25 patients with refractory AA were treated with eltrombopag using a dose escalation schedule starting at 50 mg and increasing every 2 weeks by 25 mg, if the platelet count remained less than 20x10^9/L to a maximum dose of 150 mg. The primary endpoint was hematological
response at 3–4 months. The median platelet count was 9×10^9/L (range 5–15×10^9/L), and the median TPO level in these patients was 2,767 pg/mL (range 1,615–4,618 pg/mL). It was observed that 44% of patients (11/25) demonstrated at least a response at one lineage. Of these eleven patients, four patients had bilineage response and one patient had trilineage response. In addition, 36% (9/25) had a median platelet count increment of 44×10^9/L, 24% (6/25) had a median hemoglobin increase of 44 g/L, and 63% (9/25) had a median increased neutrophil count of 1.35×10^9/L. Seven of the eleven responders continued treatment for a median of 16 months, and six achieved a trilineage response.

In the recent update on this study by Desmond et al., a second cohort, which consisted of an additional 18 patients, were added to the original cohort. The median age of these 43 patients was 44 years (range 17–77 years). These patients were heavily treated with a median of two courses of IST and transfusion dependent for red blood cells and platelets. Of these 43 patients, 40% (17/43) showed hematologic response at 3–4 months (Figure 3). Of the 14 responders who continued the treatment for a median of 16 months, seven (50%) patients achieved trilineage response. Of the five patients with near-normal blood counts who had discontinued eltrombopag at a median of 20.5 months of therapy (range 9–37 months), all continued to have stable counts for a median of 13 months (range 1–15 months). Of the 26 nonresponders, two patients demonstrated hematological improvement after discontinuing eltrombopag. A total of eight patients, two responders, and six nonresponders developed clonal evolution on eltrombopag, including five with chromosome 7 abnormalities. None of the studied patients developed acute myeloid leukemia. All the patients with chromosome 7 changes were successfully transplanted. The major toxicities were irreversible deranged hepatic transaminases.

In contrast to low TPO level in ITP, the TPO level in these patients is much higher than the normal values of 7–99 pg/mL; this study demonstrated unequivocal efficacy of eltrombopag in trilineage hematopoiesis in these patients. All patients had bone marrow biopsies done at study entry and every 3–6 months. Eltrombopag has demonstrated an increase in marrowularity in responders, and none had evidence of myelofibrosis. The marrow cellularity seen in patients treated with eltrombopag is different from that observed in patients treated with standard IST, where residual hypocellularity is frequently seen.

![Figure 3](https://www.dovepress.com/fig3.png)

**Figure 3** The clinical trials of eltrombopag in refractory AA.

**Note:** Data from Olnes et al. and Desmond et al.

**Abbreviation:** AA, aplastic anemia.
Discussion
The small molecular nature of eltrombopag, and lack of competitive binding with endogenous TPO, may maximize the potential for eltrombopag to enter the stem cell niche leading to direct expansion of the few HSCs that remain in SAA. In addition, the use of TPO-receptor agonist in patients with ITP has been postulated to increase Treg activity by increasing total circulating transforming growth factor-β1 levels. In patients with SAA, a similar increase in Treg activity with eltrombopag may provide an immunomodulatory role in the treatment of SAA.

The stimulation of HSC driven by eltrombopag may lead to genomic instability and clonal evolution to MDS or acute myeloid leukemia. The development of monosomy 7 in five patients is a concern. Monosomy 7 is a high-risk cytogenetic clone associated with an evolution to acute myeloid leukemia. It is probable that these clones exist at low levels, beyond conventional cytogenetics, with a diagnosis of SAA, and the use of targeted DNA sequencing to detect such clones may be a future direction to exclude such patients from eltrombopag.

Eltrombopag is, however, tolerated well by the majority of patients with no toxicity and absence of clonal evolution.

In view of encouraging and propitious results from Olnes et al, eltrombopag has been shown to have the potential to be effective as initial therapy. On the back of these results, both the European Medicines Agency and the US Food and Drug Administration have granted expedited approval for the use of eltrombopag in adults with acquired SAA who did not respond to IST and cannot receive HSCT. There are two further ongoing nonrandomized pilot Phase II studies evaluating its use as a first-line therapy in patients with SAA and moderate AA with significant thrombocytopenia or anemia (NCT01703169 and NCT013228587). Another two clinical trials have been started to study its safety and effectiveness by adding eltrombopag to IST in SAA (NCT02099747 and NCT01623167). One of the studies, NCT01623167, has included children with AA aged 2 years and above. These clinical trials are aimed to address unresolved doubts, such as selection of candidates, duration of therapy, safety of prolonged therapy, and risk of clonal evolution. Additional studies may also be appropriate for constitutional AA where the risk and long-term toxicities of HSCT are much higher.

Conclusion
Eltrombopag has increasingly convincing efficacy in restoring multilineage hematopoiesis. There is no doubt that it is going to change the strategy for managing AA in the future. It can be potentially used as a frontline therapy in moderate to severe AA alone or in combination with IST, as well as a salvage therapy for patients who are refractory to IST or relapse after IST. Its potential to be used as monotherapy in AA could be explained by its action on c-Mpl receptors, which present on more primitive progenitor HSCs. Moreover, the convincing results from Olnes et al and Desmond et al have highlighted its role in patients with refractory AA despite having a median of two courses of IST. The use of eltrombopag monotherapy could reduce IST treatment-related complications, such as immunosuppression, hypertension, and renal dysfunction. Results from ongoing trials will provide us robust data on its use in AA with an optimal strategy.

Disclosure
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