MISE AU POINT/IN-DEPTH REVIEW
AN UPDATE ON THALASSEMIA INTERMEDIA

Joseph E. MAAKARON¹, Maria Domenica CAPPELLINI², Ali T. TAHER³


RÉSUMÉ : La thalassémie intermédiaire englobe un groupe de maladies diverses résultant d’un déséquilibre entre la production des chaînes alfa et bêta qui aboutit à une hémolyse chronique, une érythropoïèse inefficace et une surcharge en fer. Les complications engendrées par cette maladie sont un changement dans la constitution des os, un état d’hypercoagulabilité, et des organes majeurs endommagés suite au surplus de fer. La dernière décennie a vu d’importantes avancées dans le traitement de la thalassémie. Sont examinées ici les toutes dernières thérapeutiques : la chélation du fer, la greffe de cellules souches et la thérapie génétique. L’introduction de deferasirox, un chélateur du fer administré une fois par jour par voie orale, a révolutionné la chélation du fer et s’est imposé comme étant un médicament sûr et efficace. La thérapie génétique a aussi été une innovation charnière dans les nouveaux traitements, surtout avec la découverte récente d’éléments génétiques et vecteurs vitaux qui permettent un meilleur contrôle et améliorent les résultats.

Mots-clés: thalassémie, thalassémie intermédiaire, chélation du fer, érythropoïèse inefficace, greffe de cellules souches, surcharge en fer, hémoglobinopathie, thérapie génétique, inhibiteur de Jak2


GENETICALLY SPEAKING

Thalassemia intermedia (TI) is in itself a spectrum of diseases. Mildly affected patients remain completely asymptomatic till adult life, with their only abnormality being mild anemia with hemoglobin (Hb) levels maintained between 7-10 g/dl. More severely affected patients

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ABSTRACT : Thalassemia intermedia is a genetically diverse group of diseases that is the result of an imbalance in the production of the alpha and beta chains with ensuing chronic hemolysis, ineffective erythropoiesis, and iron overload. Resulting complications include bone changes, hypercoagulability, and end-organ damage due to iron overload. This decade has witnessed major breakthroughs in the management of thalassemia. In this article, we examine these novelties in therapy including iron chelation therapy, stem cell transplant, and gene therapy. Iron chelation therapy has been revolutionized with the advent of deferasirox, a once-daily oral iron chelator, that has been shown to be safe and efficacious. Gene therapy was also at the core of this revolution with the discovery of novel gene elements and viral vectors allowing for better control and improved outcomes.

Keywords : thalassemia, thalassemia intermedia, iron chelation, ineffective erythropoiesis, hematopoietic stem cell transplant, iron overload, hemoglobinopathy, gene therapy, Jak2 inhibitor

HISTORICAL OVERVIEW

Thalassemia was first described by two American pediatricians, Thomas B. Cooley and Pearl Lee [1]. They reported a series of five children with severe anemia, growth retardation, and peculiar bone abnormalities [2-3]. It was first called “erythroblastic anemia.” The term thalassemia was later coined by George Whipple [4]. Thalassemia roughly translates to “anemia of the sea,” signifying the fact that most patients were of Mediterranean origin. Insight into the pathophysiology came when Sir David Weatherall and Dr. John Clegg and their colleagues were able to isolate and separate the thalassemiglobin chain. They deduced that the culprit was an imbalance in chain synthesis [5]. Their theory still holds till now. Beta-thalassemia is a defect in the synthesis of the beta chain of the hemoglobin leading to imbalance in the α/β ratio in the red blood cells (RBC), ineffective erythropoiesis, and a spectrum of anemia [6]. Thalassemia major (TM) is when the synthesis defect is profound leading to severe anemia requiring blood transfusion within the first year of life. On the other hand of the spectrum, we have thalassemia minor (Tm) which is completely asymptomatic and only manifests as a mild hypochromic, microcytic anemia. Sturgeon described a third group of patients with manifestations that are too severe to be called minor and too mild to be called major. He suggested the term thalassemia intermedia for this population [7].

Keywords : thalassemia, thalassemia intermedia, iron chelation, ineffective erythropoiesis, hematopoietic stem cell transplant, iron overload, hemoglobinopathy, gene therapy, Jak2 inhibitor

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present younger, between 2 and 6 years of age, and require transfusions for normal and sustained growth [8]. While Tm is the result of disruption of one of the beta chains and TM is the result of disruption of both of these chains, TI has a myriad of genotypes associated with it and is still a clinical diagnosis. It usually results from a homozygous or a compound heterozygous mutation [9]. Rarely, it can also result from only one chain being affected, making it dominantly inherited [10]. TI is milder than TM because of several reasons. It can be due to inheritance of a mild mutation, resulting in subnormal synthesis of beta chains instead of complete absence. It can also be due to coinheritance of determinants that increase gamma-chain production to stabilize the excess alpha chains. Coinheritance of alpha-thalassemia will also lead to a TI phenotype since less alpha chains will remain unpaired. One can inherit triplicated or quadruplicated alpha genotype associated with beta heterozygosity and manifest a TI phenotype. Polymorphisms that alter bone, iron, and bilirubin metabolism can also affect the clinical expression of the disease. These are called tertiary modifiers. Environmental and socioeconomic factors also play a role in the presentation. With so many factors in play (Table I), it is no surprise that TI has such a wide range of clinical presentations.

Some studies have aimed at stratifying TI patients by severity, and attempted to predict a genotype-phenotype relationship to better tailor the treatment guidelines [11-12]. Phadke and Agarwal devised a score based on the clinical presentation to classify patients. The score was composed of relevant clinical characteristics such as age at presentation, severity of the anemia, extent of growth retardation and bone marrow hyperplasia, blood transfusion requirements, and need for splenectomy [13]. This score might prove to be of great value in classifying patients and relating genotype to phenotype when developing treatment guidelines; however, more studies will have the final say in this. The diagnosis of TI remains a clinical one and can be very challenging at both ends of the spectrum.

PATHOPHYSIOLOGY

The alpha and beta chains in a normal red blood cell (RBC) should precisely match each other. In beta-thalassemia, underproduction of the beta chains causes excess, unstable alpha chains to deposit in the RBC, thus producing oxidative damage to the membrane and subsequent lysis of the cells – this is ineffective erythropoiesis [14]. This causes the bone marrow to hypertrophy, resulting in characteristic deformities of the skull and face. Cortical thinning and pathological fractures are also common [15]. The primary determinant of the anemia is the degree of ineffective erythropoiesis rather than the degree of hemolysis, which only plays a secondary role [16]. Instead, hemolysis is linked to the hypercoagulable state in TI [17] leading to silent infarcts [18] and pulmonary hypertension. The anemia and the ineffective erythropoiesis increase intestinal iron absorption leading to iron overload, which causes heart failure, endocrine abnormalities and others (Figure 1).

COMPLICATIONS

The combination of ineffective erythropoiesis, chronic hemolysis and iron overload eventually culminate in the complications we see in TI. Our first attempt at understanding the complications of TI came from comparing TI and TM patients from two centers in Lebanon and Italy. The results were remarkable in that some complications were exclusively found in TI patients [16]. A better understanding of the complications came from the OPTIMAL CARE study. This study was a cross-sectional survey of 584 TI patients from six centers throughout the Middle East and Italy. Different complications were correlated with different exposures and treatments [19]. The complications are discussed below and summarized in Table II.

A. Thrombophilia

TI patients are at a higher risk of thrombosis than TM patients. Out of 2,190 TI patients, 3.9% experienced a thrombotic event compared to only 0.9% of 6,670 TM patients [20]. The events were mainly venous. Splenectomized patients were at a higher risk of thrombosis than non-splenectomized patients. Additional risk factors for thrombosis were low hemoglobin concentration (< 9 g/dl) and transfusion-independence. Studies are still lacking on this subject but there are several proposed mechanisms. RBC remnants express negatively charged phosphatidylserine residues on their membranes which act as procoagulants, initiating thrombosis [21-22]. Activation of platelets, endothelial cells, monocytes, and depletion of

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<th>TABLE I MODIFIERS OF THALASSEMIA INTERMEDIA</th>
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<td><strong>Inheritance of modifiers that increase gamma chain production</strong></td>
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anti-thrombotic factors are also incriminated along with the cardiac, endocrine, and hepatic dysfunction [23]. Other risk factors include age [24], previous thromboembolic events, and family history.

These patients are important to identify and treat because sequelae include pulmonary hypertension with ensuing secondary heart failure [25] and silent brain abnormalities [18, 26]. Recommended therapies include aspirin or other platelet anti-aggregants in the case of thrombocytosis, or antiocoagulant agents such as low molecular weight heparin in patients with documented thrombosis or for those undergoing surgery. Blood transfusions might be considered as they dilute the circulating damaged RBCs. More studies are needed to establish a risk-assessment model that would stratify the patients according to their risks and guide a tailored therapy to this issue.

**B. Iron overload**

Iron is controlled by a negative feedback loop. Inflammation and excess plasma iron is a stimulant for the transcription of hepcidin. Hepcidin interacts with ferroportin – the iron exporter – on the basolateral surfaces of hepatocytes and the intestinal epithelium and causes its internalization and degradation [27]. This traps iron inside the cells and makes it less available in the blood stream. In the hepatocytes, macrophages, and other cells of the reticuloendothelial system, it is stored in its ferritin-bound form for later use. In the endothelial cells, intracellular iron is shed with the shedding of the epithelium. Hypoxia, anemia, and a demand for erythropoiesis suppress hepcidin, making more iron available for the bone marrow through increased intestinal absorption and better recycling of catabolic iron from the reticuloendothelial system [28]. Ineffective erythropoiesis increases expression of growth differentiation factor 15 (GDF15) and hypoxia-inducible transcription factors (HIFs) [28] which cause down-regulation of hepcidin. This increases intestinal iron absorption and depletes the macrophages of their iron stores.

![Pathophysiology of thalassemia intermedia in a nutshell.](image)

**An imbalance in the ratio of alpha to non-alpha (beta) chains leads to cell lysis and ineffective erythropoiesis. This leads to characteristic bony changes, hypercoagulability, and increased gastrointestinal iron absorption. Osteoporosis and fractures, silent brain infarcts, pulmonary hypertension, and iron overload ensue.**
After all the iron saturates the transferrin stores, it is transported as the toxic non-transferrin-bound-iron. The end result is its deposition in the parenchyma of the liver, heart, endocrine organs, and others.

Body iron levels are usually assessed by ferritin values. However, studies have shown that ferritin underestimates the total iron burden in TI [29]. For the same value of liver iron concentration, ferritin values in TI patients were significantly lower than TM patients. A proposed mechanism for this is that iron in transfused patients is preferentially distributed to the reticuloendothelial system, thus ferritin is more readily synthesized and exported [28]. This is in contrast to transfusion-independent TI patients, where low hepcidin depletes the reticuloendothelial system of its iron stores, thus ferritin will be low. Other methods of evaluating body iron stores include determination of liver iron concentration by biopsy or more recently by non-invasive techniques such as R2 MRI.

C. Splenectomy and cholecystectomy

The main indications for splenectomy in TI are poor growth and development, increased transfusion demand, hypersplenism, and splenomegaly [30]. However, splenectomy appears to contribute to an increased risk of thrombosis [22, 31]. Splenectomy is now less common than before and is performed later in life [8]. Gallstones are common in TI because of the ineffective erythropoiesis and peripheral hemolysis. That is why the gallbladder should be inspected during splenectomy to avoid further complications.

D. Extramedullary hematopoiesis

Extramedullary hematopoiesis (EMH) is when erythropoietic tissue outside marrow spaces in bones increases to compensate for the chronic state of anemia that accompanies TI. This will lead to the formation of erythropoietic masses that primarily affect the spleen, liver, and lymph nodes [8]. These masses can cause local symptoms such as neurological symptoms and spinal cord compressions [32-33]. Splenic enlargement can cause symptoms such as early satiety, left upper quadrant pain, and even raises concerns about splenic rupture [30]. Extramedullary hematopoiesis is routinely managed by radiotherapy, hydroxyurea, or transfusion therapy [32, 34-35].

E. Pulmonary hypertension

Pulmonary hypertension (PH) is defined as pulmonary artery pressure of > 25 mmHg at rest or > 30 mmHg during exercise [36]. In TI, the prevalence is estimated to be around 59%. This, in turn, contributes to an increased incidence of congestive heart failure [37]. The etiology of PH in TI is multifactorial. PH results from interplay of vasoconstriction, vascular smooth muscle proliferation, and dysfunctional endothelium with resulting thrombosis, all of which contribute to luminal narrowing and right-sided failure. Major risk factors include advancing age and a history of splenectomy [38]. Autopsy studies in thalassemia patients have revealed common histopathological findings that include plexiform and concentric medial hyperplastic pulmonary vascular lesions, and in situ pulmonary artery thrombosis [39-40]. The factors that culminate in these pathologies are most likely the result of the long-term sequelae of splenectomy, red cell membrane pathology, coagulation abnormalities, excess arginase activity, low nitric oxide bioavailability, platelet activation, oxidative stress, iron overload, and chronic hemolysis [41-45]. The correlation of ferritin with PH is still controversial with one study indicating no correlation [46] while another study reporting a strong correlation between an elevated tricuspid regurgitant jet velocity (TRV) and serum ferritin [47]. Hemolysis disables the arginine-nitric oxide pathway by releasing erythrocyte arginase [48] and cell-free hemoglobin [43]. With the consumption of nitric oxide and arginine, endothelial dysfunction ensues, and along with the intravascular hemolysis, this will create a procoagulant state [49]. Iron overload, and the subsequent oxidative stress, can also contribute to increasing the pulmonary vasculature resistance through various mechanisms. They induce pulmonary fibrosis and affect the pump itself through cardiac siderosis [50]. The hypoxemia, along with the chronic anemia, will also exacerbate the vasoconstriction [38].

Currently, there are no guidelines for the treatment of PH in TI specifically. In one small trial, PH was absent in patients taking hydroxyurea, suggesting a protective role, but further studies are needed [51]. Two small studies reported improvement in exercise tolerance in response to sildenafil [52-53]. General recommendations for the management of PH now are adequate treatment of thalassemia, any precipitating factors or associated diseases, in addition to supportive measures [38]. A multidisciplinary approach is needed and a cardiopulmonary specialist should be on board.

F. Leg ulcers, endocrine dysfunction, and hepatitis

TI patients are also at risk for other complications. Leg ulcers occur more commonly in older patients. Leg ulcers develop in some patients with low hemoglobin and do not develop in others, even though they are maintained at the same amount of fetal hemoglobin, implying that it is multifactorial in origin. The skin might be at a lower oxygen tension due to the anemia, making it fragile and more prone to break with minimal trauma. These also cause impaired healing. These are usually managed by an intense transfusion regimen to elevate the hemoglobin concentration by a minimum of 2 g/dl. Other possible interventions include elevating the end of the bed so as to keep the lower limbs at a level above the heart. This provides better venous return and increases the perfusion pressure of the affected area [8]. Zinc supplementation [54], pentoxifylline, and hydroxyurea, with or without erythropoietin [55] are also possible options.

While endocrine dysfunction can be quite common in TM, it is quite rare in TI. Patients generally experience late puberty, but their sexual development is normal and their fertility preserved. However, hypothyroidism...
Viral hepatitis is much rarer in TI than in TM due to the fact that TI patients are less often transfused. Nevertheless, the risk is not absent and careful monitoring should be undertaken [56]. New complications, such as hepatocellular carcinoma are emerging as patients are living longer.

**MANAGEMENT**

**A. Iron chelation therapy ± transfusion therapy**

The management of TI has come a long way this decade with the new oral chelators, gene therapy, and Jak2 inhibitors [Table III]. Transfusion is the only available salvage therapy when severe symptoms of anemia set in, including developmental delay in the pediatric age group. Associated risks include alloimmunisation, which is relatively common in TI, especially if transfusions are instituted after the age of 12 months [57]. Rhesus and Kell phenotyping is recommended prior to transfusion [58], with some physicians advocating a short course of steroids for 3-5 days concomitantly, even though this remains controversial.

The rate of iron loading in TI is variable and changes with transfusional load. Iron loading in non-transfused patients is estimated to be 2-5 grams per year [59] compared to 7.5-15.1 grams per year for transfused patients [60]. Iron loading should always be assessed by liver iron concentration (LIC), preferably by non-invasive MRI imaging, and iron chelation therapy initiated accordingly. Previously, an LIC of 7 mgFe/g dry weight was used to initiate therapy; however, a recent study has found that an LIC of 7 mgFe/g dry weight is associated with vascular complications and an LIC of 6 mgFe/g dry weight is associated with endocrine comorbidities, meaning that complications would have already set in, and chelation should start before that [61].

The gold standard remains desferrioxamine injections which is given subcutaneously or intravenously for prolonged periods of time and is associated with significant patient discomfort, decreased quality of life, and non-compliance [62-63] despite significant mortality and morbidity reductions [64]. Deferasirox, a once-daily oral iron chelator, has been found to be safe and efficacious in reducing iron burden in TI [65].

A novel oral iron chelator is also under development for clinical use. The molecule is still in phase II clinical trials, but the results seems promising [66]. Extensive preclinical toxicological studies demonstrated a higher no-observable-adverse-effect level (NOAEL) compared to deferasirox.

**B. Stem cell transplantation**

Hematopoietic stem-cell transplantation is essentially curative in properly selected candidates. Good candidates with high success rates include pediatric patients who have not developed complications such as viral hepatitis or severe iron overload and who receive HLA-identical related donor stem-cell transplants. The event-free survival for beta-thalassemia patients reported from several groups is 80-90% with less than 10% mortality and minimal morbidity, apart from impaired fertility [67-70]. The problem is in the availability of donors. Disease-free survival is lower, and morbidity and mortality are higher with unrelated haploidentical donors [71-73]. Definitive hematopoietic stem cells can be obtained from cord blood. Several studies have assessed the use of cord-blood transplantation for patients with thalassemia [69], and the outcomes of related cord-blood stem cell transplantation for beta-thalassemia are approaching those of conventional bone marrow transplantation with disease-free survival of 90%. A cord-blood bank will also allow unrelated transplants to take place, but data is still scarce [74-76].

The physician, patient, and everybody concerned should be involved in the transplantation decision given the associated morbidity, mortality, and cost of the procedure (Figure 2).

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**TABLE III**

**MANAGEMENT OF THALASSEMIA INTERMEDIA**

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<th>Non-conventional</th>
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<td>Transfusion therapy</td>
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<td>Iron chelation therapy</td>
<td>Splenectomy</td>
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<td>Splenectomy</td>
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**Hb**: hemoglobin
C. Gene therapy

Thalassemia is an ideal disease for gene therapy since its genetic defect only affects the erythroid lineage in the hematopoietic system. The concept is simple, albeit the mechanism is technically demanding. Hematopoietic stem cells are harvested from the patient and infected with a virus carrying the beta-globin gene along with its upstream regulatory elements, and then infused back after myelosuppression. The fact that the patient’s own cells are used overcomes the problem of scarcity of donors and associated immunological complications.

A lot of work has been put to develop viral constructs that are able to accommodate all the sequences needed for the expression of the beta-globin gene. “Boundary elements” are now frequently included in the genetic complexes to include proper insertion. Boundary elements are sequences that flank the gene and delimit the activity of the regulatory elements of that gene [77]. Another setback was the associated risk of leukemia with the use of retroviral vectors. Therefore, lentiviral vectors were developed and they have proven successful in mouse models both in TM and T1 [78-83]. A clinical trial to test this lentiviral vector has already begun in 2007 on two patients [84-85]. One of the patients needed rescue with non-manipulated cells while the other achieved 10% engraftment and was able to become transfusion independent. The future of gene therapy probably lies in induced pluripotent stem cells, which are human somatic cells that are reprogrammed to form multipotent stem cells [86-87].

D. Jak2 inhibitors

Data showed that in ineffective erythropoiesis, there is an accumulation of proliferating erythroid progenitors, and an increase in the percentage of erythroid cells in the S-phase. The thalassemia cells were associated with the expression of certain cell-cycle regulatory genes such as EpoR, Jak2, cyclin-A, Cdk2, Ki-67 and the antiapoptotic protein Bcl-XL, and differentiated less. In thalassemic mice, a Jak2 inhibitor reversed ineffective erythropoiesis and decreased spleen size with limited effect on anemia. Jak2 inhibitors would be expected to limit or reduce splenomegaly, thus improving anemia and delaying the need for splenectomy. A decrease in ineffective erythropoiesis is also expected to upregulate hepcidin, thus improving the metabolism of total body iron content and decreasing end-organ damage [88-90].

CONCLUSION

Despite all the advances in premarital screening and genetic counseling, thalassemia remains an important genetic disease. With migratory patterns and intermixing of populations, thalassemia is no longer limited to its places of origin. This decade has witnessed breakthroughs in the treatment of thalassemia and we are now able to provide better medical care that is making thalassemia more of a chronic disease than a fatal one.


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