Iron overload and chelation therapy in hemoglobinopathies

Rayan Bou-Fakhredin, Joseph Elias, Ali T. Taher

Department of Internal Medicine, American University of Beirut Medical Center, Beirut, Lebanon

Abstract

Iron overload (IOL) is highly prevalent among patients with hemoglobinopathies; both transfusion dependent thalassemia (TDT) and non-transfusion dependent thalassemia (NTDT). Whether IOL is secondary to regular transfusions like in TDT, or develops from increased intestinal absorption like in NTDT, it can cause significant morbidity and mortality. In TDT patients, iron accumulation in organ tissues is highly evident, and leads to organ toxicity and dysfunction. IOL in NTDT patients is cumulative with advancing age, and concern with secondary morbidity starts beyond the age of 10 years, as shown by the OPTIMAL CARE study. Several modalities are available for the diagnosis and monitoring of IOL. Serum ferritin (SF) assessment is widely available and heavily relied on in resource-poor countries. Non-invasive iron monitoring using MRI has become the gold standard to diagnose IOL. Three iron chelators are currently available for the treatment of IOL: deferoxamine (DFO) in subcutaneous or intravenous injection, oral deferoxiprone (DFP) in tablet or solution form, and oral deferasirox (DFX) in dispersible tablet (DT) and film-coated tablet (FCT). Today, the goal of ICT is to maintain safe levels of body iron at all times. Appropriate tailoring ICT with chelator choices and dose adjustment must be implemented in a timely manner. Clinical decision to initiate, adjust and stop ICT is based on SF, MRI-LIC and cardiac T2*. In this article, we review the mechanism of IOL in both TDT and NTDT, the pathophysiology behind it, its complications, and the different ways to assess and quantify it. We will also discuss the different ICT modalities available, and the emergence of novel therapies.

Introduction

Thalassemia is an inherited disease with multiple genetic forms, including alpha-thalassemia, beta-thalassemia, hemoglobin E/beta thalassemia, and others. Molecular defects in the alpha-globin gene cluster on chromosome 16 or the beta-globin gene cluster on chromosome 11 result in defective hemoglobin synthesis. This in turn leads to an imbalance in the relative quantity of alpha-globin and beta-globin chains [1]. Therefore, the disease hallmarks consist first and foremost of the before mentioned imbalance in the α/β-globin chain ratio, which in turn leads to the following cascade of events and disease hallmarks including ineffective erythropoiesis, and chronic hemolytic anemia (Figure 1).

Thalassemic disorders lie on a spectrum of severity with different clinical phenotypes, complications, and strategies for treatment. The grade of this severity relies on the significance of the globin gene mutation and coinheritance of other genetic determinants. [2] The degree of transfusion dependence is one of the elements considered in a recent classification of thalassemic disorders into transfusion-dependent thalassemia (TDT) and non–transfusion-dependent thalassemia (NTDT). Iron overload (IOL) is highly prevalent among patients with hemoglobinopathies. Whether IOL is secondary to regular transfusions like in TDT, or develops from increased intestinal absorption like in NTDT, it can cause significant morbidity and mortality. In TDT patients, iron accumulation in organ tissues is highly evident, and leads to organ toxicity and dysfunction. In NTDT patients, IOL is cumulative with advancing age, and concern with secondary morbidity starts beyond the age of 10 years. [1] In this article, we review the mechanism of IOL in both TDT and NTDT, the pathophysiology behind it, its complications, and the different ways to assess and quantify it. We will also be addressing the different ICT modalities available, and discuss the emergence of novel therapies targeting IOL.

Mechanism of iron overload in transfusion dependent thalassemia and non-transfusion dependent thalassemia

The predominant mechanisms driving the process of iron loading include increased iron burden secondary to transfusion therapy in TDT and enhanced intestinal absorption secondary to ineffective erythropoiesis and hepcidin suppression in NTDT. Different organs are affected differently by iron overload in TDT and NTDT owing to the underlying iron loading mechanism and rate of iron accumulation[1].

Unfortunately, the human body lacks a physiological mechanism for removal of the excess iron load resulting from blood transfusion[3]. Each unit of transfused packed red blood cells contains 200 to 250 mg elemental iron. In TDT, transfusional iron usually amounts to 0.3 to 0.6 mg/kg per day with an assumed monthly transfusion rate of 2 to 4 U packed red blood cells. Senescent transfused red blood cells are phagocytosed by the reticuloendothelial macrophages. This leads to the release of cellular iron into the...
plasma to bind transferring; which is the main iron transport protein and is capable of binding two Fe^{3+} Molecules at once. It is only when transferring binding gets saturated, that we start having iron accumulation: because the now non-transferrin-bound iron (NTBI) is readily transported through calcium channels into the liver (hepatocytes), heart (cardiac myocytes), and endocrine glands. The accumulation of iron in different organs leads to the different clinical complications of IOL [1, 3]. This accumulation of NTBI in different types of cells leads to its metabolism and the production of reactive oxygen species (ROS) contributing to the cellular dysfunction, apoptosis, and necrosis seen in the target organs[3, 4].

Transferrin carrying 2 molecules of Fe^{3+} then binds to transferrin receptor 1 (TfR1) and transferrin receptor 2 (TfR2), then gets endocytosed. The acidic environment of the lysosomes, causes the release of Fe^{3+} from transferrin, and its reduction to Fe^{2+}. Fe^{2+} then reaches the cytosol through divalent metallic transporter 1[1]. While TfR2 is uniquely expressed in the liver and intestine, TfR1 is expressed in most tissues, including erythroid precursors, the liver, and the myocardium. The affinity of TfR1 for iron is higher than that of TfR2 by ~25 times. Interestingly, TfR2 lacks an iron responsive element, and iron loading continues to happen in the liver despite high liver iron concentration (LIC), while TfR1 is downregulated with elevated transferrin saturation. Previously, the most important clinical complication of iron overload was cardiac siderosis, which is at the origin of arrhythmias and heart failure and has been a major cause of mortality in TDT. However, with the advances in IOL diagnosis and management nowadays, cardiac mortality has declined significantly, allowing light to be shed on hepatic and endocrine dysfunction as other complications of IOL in TDT patients[4].

Even in the absence of regular red blood cells transfusions, IOL still develops in patients with NTDT. Remarkably, it has been noted that iron accumulation preferentially occurs in the liver in patients with NTDT and rather than the myocardium. This was established after observational studies showed absence of cardiac siderosis even in patients with severely elevated liver iron content (LIC) [5]. Normally, Hepcidin synthesis by the liver suppresses the release of iron from erythroid precursors, hepatocytes, basolateral membranes of hepatocytes, and macrophages by binding to ferroportin, which mediates iron export[1]. It is believed that the ineffective erythropoiesis, along with the state of chronic anemia/hypoxia leads to the inappropriately low levels of hepcidin. This in turn contributes to IOL through two mechanisms: increased intestinal iron absorption through lowering ferroportin, and increased release of recycled iron from the reticuloendothelial system. This in turn leads to preferential portal and subsequently hepatocyte iron loading, depletion of macrophage iron, and relatively lower levels of serum ferritin (compared to TDT patients) [6].

Iron overload complications in transfusion dependent thalassemia and non-transfusion dependent thalassemia

Iron overload complications in transfusion dependent thalassemia

As previously mentioned, when iron content surpasses transferrin binding capacity and hemosiderin/ferritin storage ability, toxic NTBI enters mitochondria and leads to the formation of toxic radicals and ROS. This in turn leads to gene alterations resulting in cell apoptosis and or fibrosis in different target organs including the myocardium, liver, and endocrine glands [7]. Given the constant need for blood transfusions in TDT, along with hypoxia/anemia induced hepcidin suppression, IOL occurs at a faster rate in TDT patients when compared to NTDT patients. This is also evident in the clinical course of the two diseases, as IOL complications are more pronounced in the TDT group.

a. Cardiac complications – Despite the advances in ICT, cardiovascular disorders remain the leading cause of morbidity in TDT patients, and it remains crucially important for clinicians to recognize it early on as it mandates intensified chelation therapy. IOL related cardiac complications include reversible myocyte injury, arrhythmias including heart block, and arterial changes with loss of vascular compliance [7-9].

b. Hepatic complications – The liver is another organ that is highly susceptible to IOL induced damage in TDT patients. Hepatic macrophages known as Kupffer cells are primarily affected due to their role in RBC degradation. The intra-macrophagic iron will be released in the bloodstream in a progressive manner. During this process, plasma transferrin gets saturated which leads to the appearance of NTBI. NTBI will in turn target different organs including the heart, liver and endocrine organs. Hence controlling liver iron content is crucial to protect the liver along with the other organ systems [10]. Hepatic complications comprise hepatic fibrosis, cirrhosis, and in cases hepatocellular carcinoma [11].

c. Endocrine and bone complications – Endocrine disorders seen in TDT patients due to IOL include short stature and growth retardation, hypogonadism and delayed puberty, hypothyroidism, impaired glucose tolerance and diabetes mellitus, hypoparathyroidism, adrenal insufficiency as well as osteoporosis [12, 13].

Iron overload complications in non-transfusion dependent thalassemia

NTDT is associated with a high morbidity profile that can start manifesting as early as 10 years of age. Iron burden and accumulation stands behind some of the complications seen in NTDT: by promoting oxidative damage in different organs of the body, and inducing multiple organ dysfunction.

a. Cardiac Disease – Whereas cardiac disease is one of the most important causes of morbidity/mortality in patients with TDT,
it is manifested in a less severe manner in patients with NTDT. In fact, while IOL in TDT leads to left ventricular (LV) dysfunction, heart failure and in severe cases to cardiogenic shock, cardiac complications in NTDT are related to right sided heart failure secondary to pulmonary hypertension [8]. However, the risk of LV decompensation in patients with NTDT is minimal, but increases with age. This might be explained by the fact that iron deposition in the myocardium happens at a faster rate and is much more common in patients with TDT when compared to patients with NTDT [14]. Moreover, when compared to healthy individuals, patients with NTDT were found to have a higher prevalence of rhythm disorders, pericardial diseases and valvular abnormalities [8, 9].

b. Liver Complications – In NTDT, most iron accumulation targets the liver, and patients are at an increased risk of hepatic fibrosis, cirrhosis, and eventually developing hepatocellular carcinoma (HCC). IOL is one of the most important risk factors for hepatic failure and cirrhosis seen in-thalassemia patients. Even in the absence of chronic hepatitis C (HCV) infection, NTDT patients are at an increased risk of HCC due to IOL. As already mentioned, iron accumulation is associated with the formation of free radicals and reactive oxygen species (ROS) in different target organs, specifically hepatocytes leading to cellular damage by inflicting damage to tumor suppressor genes and DNA repair genes. Moreover, iron has a pro-biogenic effect which in turn accelerates the development of liver cirrhosis [15, 16].

c. Endocrinopathies – Iron accumulation can also disrupt the hypothalamic-pituitary axis (HPA), leading to an array of endocrine diseases in-thalassemia patients such as hypogonadism, hypothryroidism, hypoparathyroidism, diabetes mellitus, and adrenal insufficiency. Endocrinopathies form an important source of morbidity in NTDT patients, but with a lower prevalence than their counterparts in TDT. This may be attributed to the hepatic dominance of iron loading and to the slower rate at which IOL occurs in patients with NTDT [17, 18].

d. Kidney disease – Kidneys are also affected by iron overload, manifesting as proteinuria, and glomerular hyperfiltration as a result of glomerular and tubulointerstitial injury. It is important to mention that damage to the glomerulus and the tubulointerstitial systems are not only caused by IOL; the chronic state of anemia and hypoxia play a crucial role in this too. End stage kidney disease is a possible outcome of IOL induced renal damage in patients with NTDT [19, 20].

e. Bone Disease – Iron overload, splenectomy, low fetal hemoglobin levels and female gender appear to be associated with a higher risk of osteoporosis in NTDT patients [21-23]. On the other hand, ICT and hydroxyurea use were correlated with lower rates of osteoporosis.

**Magnetic resonance imaging**

Given its safety and reliability when compared to the invasive liver biopsy, MRI using T2* (in milliseconds) and R2* imaging techniques are now considered the gold standard for LIC quantification. T2* relaxation refers to decay of transverse magnetization caused by a combination of spin-spin relaxation and magnetic field inhomogeneity[24]. This relaxation occurs faster (shorter T2* in ms) with increasing tissue (myocardial, liver…) iron concentrations [1, 24].

Moreover, Angelucci et.al demonstrated that LIC estimated from MRI imaging in mg of iron per gram of liver dw correlates reliably with total body iron stores [10]. Most guidelines now rely on LIC and T2* from MRI to diagnose IOL and adapt ICT [1]. Specific LIC and cardiac T2* thresholds have been associated with morbidity in TDT and NTDT [11, 25]:

- In NTDT, LIC values greater than 5 mg/g dw were associated with increased morbidity.
- In TDT, LIC values greater than 7 mg/g dw are used to indicate increased risk for complications related to iron overload, while LIC values >15 mg/g were predictive of advanced liver fibrosis, mortality, and increased risk of cardiac disease in TDT [10, 25].
- Cardiac T2* values lower than 10 ms is highly associated with increased risk of symptomatic heart failure and higher mortality in TDT [26].
- Cardiac T2* values between 10 and 20 ms were associated with lower left ventricular ejection fraction (LVEF) and a higher risk of arrhythmias in TDT [26].

**Serum ferritin assessment**

The unavailability of MRI in the developing countries where Thalassemia is most prevalent (Sub-Saharan Africa, Middle East, India, Mediterranean region, and Southeast Asia), and its high cost form major limitations to its use as a tool to quantify and guide ICT. Therefore, the assessment of serum ferritin values and their correlation with LIC becomes of crucial value. In NTDT for example, cutoffs of 300 ng/mL and 800 ng/mL were identified: whereas SF levels lower than 300 ng/mL indicate absent IOL and SF levels higher than 800 ng/mL indicate significant IOL [27]. Moreover, results from the ORIENT study revealed that patients with SF ≥800 μg/L have a higher incidence of morbidities over 11 years. Based on a ROC analysis, a SF level of ≥800 μg/L had the highest accuracy for predicting LIC ≥5 mg Fe/g dw. As for the SF values between 300 and 800, a recent evaluation found that a significant proportion of those NTDT patients had IOL requiring treatment [28]. As for TDT, the cutoff used to initiate ICT is 1000 ng/mL. Since SF levels lower than 1000 ng/mL were associated with lower morbidity and mortality in TDT and this threshold is most commonly used to indicate the need for initiation and as a target for [4, 29]. In a multicenter study conducted in 2017 by Krittayaphong et al. SF levels had limited ability to correlate with cardiac iron overload in TDT, but were found to predict cardiac siderosis when values were greater than 2500 ng/mL [30].

**SQUID**

LIC can also be derived from the paramagnetism in the liver. This can be measured using superconducting quantum interference device known as SQUID [31]. However, three major drawbacks made SQUID a rarely used technique for LIC measurement: it is costly since it utilizes liquid helium, the apparatus needs to be away from all paramagnetic forces (cars, lifts…) making it impractical, finally it relies on strong mathematical methods and different calibration methods between different devices making the comparison of results between different devices trickier [31].

**Iron chelation therapy**

Iron chelation therapy (ICT) is and will always remain the standard method of choice in-thalassemia management, decreasing morbidity and mortality in this patient population. The primary goal of ICT today has shifted from treating or rescuing IOL to maintaining safe levels of body iron at all times [32]. Moreover, appropriate tailoring ICT with chelator choices and dose adjust-
Iron chelation therapy in transfusion dependent thalassemia

In TDT patients, choices of ICT monotherapy may vary. As first line of treatment the following is recommended: DFO 30-60 mg/kg/day, administered over a span of 8–10 hours a day, 5–7 days a week; or DFX 20-40 mg/kg/day administered once daily [32]. For second line treatment, and when ICT with DFX or DFO is inadequate, DFP is given at a dose of 75-100 mg/kg/day divided over three doses [32]. Possible combination therapies that have been recommended include DFO+DFP, DFO+DFX and DFP+DFX [35-37]. Indications to intensify ICT in TDT: SF ≥2500 ng/mL and/or LIC >7 mg/g dry wt. liver and/or cardiac T2* <20 msec. In TDT patients, the indication to stop ICT: SF <300 ng/mL and/or LIC <3 mg/g dry wt. liver [32] (Table 2).

Table 1. Characteristics of the currently available iron chelators in thalassemia management [2, 77].

<table>
<thead>
<tr>
<th></th>
<th>DFO</th>
<th>DFP</th>
<th>DFX</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structure</td>
<td><img src="image" alt="DFO" /></td>
<td><img src="image" alt="DFP" /></td>
<td><img src="image" alt="DFX" /></td>
</tr>
<tr>
<td>Administration route</td>
<td>Subcutaneous or Intravenous</td>
<td>Oral (tablets or solution)</td>
<td>Oral (dispersible tablet or film-coated tablet)</td>
</tr>
<tr>
<td>Administration time</td>
<td>Every 8-12 hours ± 7 days/week</td>
<td>3 times daily</td>
<td>Once daily</td>
</tr>
<tr>
<td>Half-life</td>
<td>20-30 minutes</td>
<td>3-4 hours</td>
<td>12-16 hours</td>
</tr>
<tr>
<td>Recommended dose</td>
<td>30-60 mg/kg/day</td>
<td>75-100 mg/kg/day</td>
<td>TDT: 20-40 mg/kg/day NTDT: 5-20 mg/kg/day</td>
</tr>
<tr>
<td>Route of iron excretion</td>
<td>Urinary and fecal</td>
<td>Urinary</td>
<td>Fecal</td>
</tr>
<tr>
<td>Adverse Events</td>
<td>Delay in bone growth, auditory and ocular complications, local reactions and allergies</td>
<td>Gastrointestinal complications, Neutropenia/agranulocytosis, arthralgia, elevated hepatic enzymes</td>
<td>Gastrointestinal bleeding ulceration, and irritation, elevated hepatic enzymes, increased creatinine, liver failure and renal insufficiency, skin rashes</td>
</tr>
</tbody>
</table>

TDT, Transfusion-dependent thalassemia; NTDT, non-transfusion-dependent thalassemia; DFO, deferoxamine; DFP, deferiprone; DFX, deferasirox.

Table 2. Iron overload characteristics and iron chelation therapy indications in transfusion dependent thalassemia vs non-transfusion dependent thalassemia [78].

<table>
<thead>
<tr>
<th></th>
<th>TDT</th>
<th>NTDT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanism of IOL</td>
<td>Blood transfusion</td>
<td>Increased intestinal absorption</td>
</tr>
<tr>
<td>Rate of iron accumulation</td>
<td>Fast</td>
<td>Slow</td>
</tr>
<tr>
<td>IOL-related complications</td>
<td>Cardiac siderosis, heart failure and cardiac arrhythmia, liver fibrosis and cirrhosis, endocrinopathies</td>
<td>Liver fibrosis, cirrhosis and HCC, endocrinopathies, proteinuria and glomerular hyperfiltration, thrombosis, PHT, osteoporosis, osteopenia</td>
</tr>
<tr>
<td>Indication to initiate ICT</td>
<td>SF ≥1000 ng/mL or LIC ≥3 mg/g dry weight liver</td>
<td>SF ≥800 ng/mL and/or LIC ≥5 mg/g dry weight liver</td>
</tr>
<tr>
<td>Indication to intensify ICT</td>
<td>SF ≥2500 ng/mL and/or LIC &gt;7 mg/g dry wt. liver and/or Cardiac T2* &lt;20 msec.</td>
<td>LIC after 6 months of treatment &gt;7 mg/g dry wt. liver or SF &gt;1500-2000 ng/mL</td>
</tr>
<tr>
<td>Indication to stop ICT</td>
<td>SF &lt;300 ng/mL and/or LIC &lt;3 mg/g dry wt. liver</td>
<td>SF &lt;300 ng/mL and/or LIC &lt;3 mg/g dry wt. liver</td>
</tr>
<tr>
<td>Choices of ICT-Monotherapy</td>
<td>DFO 30-60 mg/kg/day</td>
<td>DFX 20-40 mg/kg/day</td>
</tr>
<tr>
<td></td>
<td>DFX 75-100 mg/kg/day</td>
<td>DFP 5-20 mg/kg/day</td>
</tr>
<tr>
<td>Choices of ICT-Combination Therapy</td>
<td>DFO+DFP</td>
<td>DFO+DFX</td>
</tr>
<tr>
<td></td>
<td>DFP+DFX</td>
<td></td>
</tr>
</tbody>
</table>

TDT, transfusion-dependent thalassemia; NTDT, non-transfusion-dependent thalassemia; SF, serum ferritin; LIC, liver iron concentration; HCC, hepatocellular carcinoma; PHT, pulmonary hypertension; DFO, deferoxamine; DFP, deferiprone; DFX, deferasirox.
(T2* 8–20 ms) showed that at 12 months, myocardial T2* improved significantly in both groups [38]. However, the between-group difference was significantly in favor of combination therapy. In comparison with the standard IOL using DFO, combination therapy with DFP showed greater reduction in myocardial iron and improved LVEF [38]. Therefore, combination therapy should be considered in patients whose DFO monotherapy fails to achieve adequate control of myocardial iron [38]. However, addition of DFP to DFO did not always enhance myocardial iron removal in patients with severe myocardial IOL. The cardiac sub study of the EPIC trial, a 1-year, multicenter, prospective longitudinal study, looked at the effect of DFX in removing cardiac iron in patients with β-thalassaemia and myocardial siderosis over 3 years and showed that DFX continually improved myocardial T2* across 3 years irrespective of baseline myocardial iron severity [39]. Different chelators have different strengths in removing NTBI and lowering mitochondrial oxidative stress. Nevertheless, DFX and DFP both improve endothelial function over time [7].

b. Approach to hepatic IOL – With increasing LIC, liver enzymes increase, as well as liver fibrosis. Organ damage and dysfunction can progress [40]. LIC is the best measure of total body iron stores and helps to predict the risk of hepatic and extrahepatic complications [41, 42]. Effective ICT reduces LIC and may prevent progression of liver disease. LIC levels >7 mg/g dw are associated with increased risk of morbidity and liver disease. LIC levels >15 mg/g dw are associated with increased aminotransferase enzymes concentration, increased risk of hepatic fibrosis and hepatocellular carcinoma (HCC), increased risk of HCC secondary to cirrhosis and increased risk of cardiac disease [40, 42, 43]. DFO therapy has been shown to be associated with a significant decrease in LIC in patients TDT. DFX has also been shown to significantly decrease LIC by 3.1 to 7.8 mg/g dw in patients with TDT [44]. One study showed that DFX doses >30 mg/kg per day were needed to achieve optimal improvement in LIC in patients with heavy IOL in TDT [45]. DFX monotherapy and DFP monotherapy have been shown to improve hepatic siderosis [35, 46, 47] However, there is no head-to-head trial comparing DFX and DFP at optimal doses. Based on the most robust data available, we recommend monotherapy with DFX, at a dosage of 20 to 30 mg/kg or higher per day, or combination therapy with DFO and DFX for the treatment of hepatic siderosis in TDT [37, 44, 48]. In patients with high LIC, combination therapy of DFX+DFO has been shown to significantly reduce LIC.

c. Approach to IOL in the endocrine organs – Endocrinopathies still account for significant morbidity in TDT [12, 49-51]. For example, pancreatic iron loading (defined as MRI R2* >100Hz) and severe pituitary iron deposition may possibly develop during the first decade of life [52, 53]. Intensive combination therapy of DFO and DFP has been associated with prevention and/or reversal of endocrine complications in general [54]. Combined DFP/DFX treatment has also been shown to prevent or reverse endocrine complications in TDT patients. In a recent multicenter retrospective cohort study by Casale et al., the low prevalence of new endocrine disorders and stabilization of preexisting ones during DFX therapy, suggest that DFX may play a protective in endocrinopathies [55].

Iron chelation therapy in non-transfusion dependent thalassemia

While all three iron chelators have proven their effectiveness as iron chelators in TDT patients, DFX remains the only drug that has received Food and Drug Administration (FDA) and European Medicines Agency (EMA) approval for use in NTDT patients, mostly based on results extracted and published from the THALassa trial [56, 57]. In this multinational, prospective, randomized, double-blinded phase II trial 1-year DFX treatment of NTDT patients >10 years was found to decrease LIC at a daily dose of 5 and 10 mg/kg, respectively, compared to placebo [57]. Sub-analyses further proved DFX 5 and 10 mg/kg/day starting doses led to consistent reductions in LIC across all patients, irrespective of baseline LIC, SF, underlying NTDT form, splenectomy status or demographics such as age, gender and race [57]. The analyses also showed that greater reductions in LIC were achieved in patients dose-escalated at 6 months from DFX 10 mg/kg/day starting dose to 20 mg/kg/day [57]. A 1-year extension phase was then carried out to allow for the assessment of up to 2 years of DFX treatment. Patients continued to respond, with a decrease in LIC and SF over 2 years. Data extracted from the THETIS study [58] further showed that a starting dose of 10 mg/kg/day of DFX is effective in reducing IOL in NTDT, and that dose escalation up to 30 mg/kg/day should be considered starting at week 4 based on LIC response [58]. DFP has not been extensively studied in NTDT. Single-arm, open-label studies with small sample sizes and a more recent randomized controlled trial showed significant decreases in SF and LIC with DFP therapy [59]. DFO has not been systematically studied in NTDT, although studies with small sample sizes and short durations have shown an increase in urinary excretion of iron and a decrease in SF.

In NTDT, specific indications have been established for the initiation, dose escalation and termination of ICT. DFX chelation with initial starting dose of 10 mg/kg/day should be started in patients ≥10 years of age (15 years of age in hemoglobin H disease) if their LIC ≥5 mg Fe/g dry weight, or if their SF concentration was found to be ≥800 μg/L when LIC is not available due to lack of the necessary MRI technology [33]. As for monitoring of iron levels, LIC should be checked 6 months after therapy initiation, with follow up every 6–12 months, in addition to SF levels being measured every 3 months [33]. If at 6 months LIC is still >7 mg Fe/g dry weight (or SF >1500 μg/L only if LIC is unavailable) with less than 15% reduction in baseline values, dose escalation should be considered up to 20 mg/kg/day [33]. DFX therapy can be safely discontinued when patients reach an LIC value of 3 mg Fe/g dry weight (or SF level of 300 μg/L only if LIC is unavailable) [33]. In NTDT, it is recommended to intensify ICT if the LIC after 6 months of treatment >7 mg/g dw. liver or SF >1500–2000 ng/mL and <15% decrease from baseline. Indications to stop ICT in NTDT include a SF <300 ng/mL and/or LIC <3 mg/g dry wt. liver (Table 2).

Adherence and advances in iron chelation therapy

Compliance with ICT is associated with effective control of IOL and improved patient survival [60, 61]. Moreover, adherence to long-term ICT is crucial in preventing IOL-related complications. For example, barriers to optimal adherence to DFX-DT include preparation time, palatability, the need to take the drug in a fasting state, and drug-related side effects, notably gastrointestinal (GI) tolerability [62]. A new FCT formulation was developed, which is swallowed once-daily, whole or crushed, with or without a light meal [62]. The open-label, phase II ECLIPSE study evaluated the overall safety, as measured by the frequency and severity of adverse events (AEs) and changes in laboratory values, in patients treated with DFX-FCT or DFX-DT. Overall incidence of AEs was similar between treatments, but there were fewer serious adverse events (SAEs) with FCT. The study also evaluated patient-reported outcomes (PRO) in TDT or lower-risk myelodysplastic
syndromes patients randomized to receive DFX DT or FCT over a 24-week period [62]. FCT recipients consistently reported better adherence, greater satisfaction, and fewer concerns, with a safety profile consistent with the known DT formulation. These findings suggest a preference in favor of the new formulation, with better patient satisfaction and adherence translating into reduced IOL-related complications.

Iron chelation therapy in special populations:
Pediatric population

Pediatric TDT patients require adequate blood transfusions for normal growth and skeletal development. The goals of blood transfusion therapy in children with TDT include correction of anemia, suppression of erythropoiesis and bone changes, prevention of spleen enlargement and hypersplenism and ensure appropriate growth and development. TDT patients may have liver IOL as young as age 2 years. Young TDT patients may also have myocardial IOL [63]. Guidelines for pediatric patients include maintaining an average Hb of 12 g/dL, max 14 g/dL, a pre-transfusion Hb of 9–10.5 g/dL, and transfusing pRBC units only: starting with low transfusion frequency in young patients and increase as they grow. Chelation strategies initiated timely and adjusted appropriately in children must be warranted to prevent permanent organ damage that might lead to significant morbidity later in life [64]. DFO has been shown to reduce serum ferritin in children with IOL [64]. However, high DFO doses in pediatric patients with low serum ferritin may result in growth failure, which should be distinguished from growth retardation secondary to inadequate transfusion or IOL [65]. In such cases, close monitoring of growth rate, DFO dose/regimen is recommended. Dose reduction was also found to restore growth rate to pre-treatment levels in some cases. Therefore, it is recommended to check on body weight and height every 3 months in children [32]. The experience in DFP is the most limited, with studies available showing it may not be as effective to reduce iron in young children [66]. DFX therapy has been shown to have a long-term efficacy in TDT patients as young as 2 years of age, with no observed negative effects on growth or sexual development [67]. It is important to mention that challenges in the treatment of thalassemia change with age. In early childhood, the clinician must ensure adequate support and therapy to optimize growth and development. In late childhood and adolescence, sexual development and transition of care are important areas of focus. As patients transition in to adulthood, the goals of therapy include preventing long-term complications related to anemia, IOL, and hypercoagulability [68].

Novel therapies targeting iron overload

Newly emerging therapies targeting iron dysregulation include minihepcidins, and transmembrane protease serine 6 (TMPRSS6) (Figure 1).

Minihepcidins

Minihepcidins, or long-acting hepcidin analogs, have been shown to restrict iron absorption, and their utilization has shown beneficial effects on ineffective erythropoiesis and consequently IOL [69-71]. These long acting hepcidin analogs have shown to increase the levels of endogenous hepcidin, thus decreasing iron absorption from the GI tract, and increasing the redistribution of iron, thereby limiting end-organ toxicity [72]. Studies conducted on mice have also shown that minihepcidin therapy can increase Hb concentrations, and also decrease reticulocyte counts in addition to reducing spleen size [72, 73].

TMPRSS6

Several studies have been reported on the use of transmembrane protease serine 6 (TMPRSS6) as an approach to stimulate endogenous hepcidin production [74-76]. TMPRSS6, a transmembrane serine protease, acts by reducing the production of hepcidin. Thus, endogenous hepcidin production can be stimulated by reducing the expression of TMPRSS6. Through data from mouse models, it has been shown that TMPRSS6 gene deletion not only improves anemia but also reduces ineffective erythropoiesis, splenomegaly, and IOL [74]. Other studies have shown that the use of antisense oligonucleotides or small interfering RNAs that target TMPRSS6 lead to improvements in anemia and IOL [75, 76]. Genetic ablation of TMPRSS6 also improved ineffective erythropoiesis and decreased splenomegaly in NTDT patients, without a concomitant decrease in erythropoietin production [74].

Conclusions

In conclusion, IOL represents an important clinical problem in thalassemia patients. Adequate assessment and monitoring of IOL in TDT and NTDT patients, in addition to tailored ICT, is crucial for preventing the complications known to be associated with this increased iron burden. New treatment modalities are currently being investigated to broaden options available for TDT and NTDT management, with ultimate goals of prolonging longevity, promoting greater compliance and better adherence and improving quality of life. Since both TDT and NTDT patients present with multiple pathophysiologies, tailoring treatment will always remain essential.

References

9. Amoozgar H, Zeighami S, Haghpanah S, Karimi M. A comparison of heart function and arrhythmia in clinically asymptomatic patients with beta thalassaemia intermedia and beta-tha-
24. Chavhan GB, Babyn PS, Thomas B, Shroff MM, Haacke EM.

22. Musallam KM, Taher AT, Cappellini MD, Sankaran VG.

20. Mallat NS, Musallam KM, Mallat SG, Ziyadeh FN, Koussa S,

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,

13. Toumba M, Skordis N. Osteoporosis syndrome in thalassaemia

10. Angelucci E, Brittenham GM, McLaren CE, Ripalti M, 

Baronciani D, Giardini C, et al. Hepatic iron concentration and 

16. Kowdley KV. Iron, hemochromatosis, and hepatocellular car-

15. Kew MC. Hepatic iron overload and hepatocellular carcinoma.

14. Taher AT, Musallam KM, El-Beshlawy A, Karimi M, Daar S,


