INTRODUCTION

The hemoglobinopathies constitute the most common monogenic disorders in humans. Among them, thalassemia results from defective globin chain synthesis leading to chronic hemolytic anemia. Although highly prevalent and once confined to certain geographic areas, including the Mediterranean Basin, Middle East and Southeast Asia, thalassemia now has a global distribution owing to the migration of populations from high-prevalent regions to the Western World, a phenomenon that has lately been intensified due to the massive migration movements from Middle East and Asia to Europe. Thalassemia and particularly transfusion-dependent thalassemia (TDT) is a demanding clinical condition, requiring lifelong care and follow-up, ideally in specialized centers and by multidisciplinary teams of experts, and results in huge healthcare expenditures. Although advances in diagnosis and treatment have dramatically improved prognosis and survival of patients with thalassemia, offering them the possibility of a nearly normal lifespan with high quality, these advances have not been equally adopted and applied to all patient populations and this is particularly true in some highly prevalent areas. In addition, the healthcare systems in western countries, where the disease has always been extremely rare, are not ready to address the special needs of migrant thalassemia populations. Therefore, the management of patients with thalassemia remains challenging.

Thalassemia International Federation (TIF) is a global patient-driven umbrella federation with 232 member-associations in 62 countries across the world, operating in official relations with the World Health Organization, the United Nations Economic and Social Council and the European Commission. TIF aims at ensuring equal access to quality care for all thalassemia and other hemoglobinopathy patients in every part of the world by promoting education, research, awareness, and advocacy. One of TIF’s main actions is the development and dissemination of clinical practice guidelines for the management of these patients. In 2021, the fourth edition of TIF’s guidelines for the management of TDT was published. The full text provides detailed information on the management of TDT patients and the clinical presentation, pathophysiology, diagnostic approach, and treatment of disease complications or other clinical entities that may occur in these patients, while also covering relevant psychosocial and organizational issues. The present document is a summary of the 2021 TIF guidelines for TDT that focuses mainly on clinical practice issues and recommendations.

ABSTRACT

Beta-thalassemia and particularly its transfusion-dependent form (TDT) is a demanding clinical condition, requiring life-long care and follow-up, ideally in specialized centers and by multidisciplinary teams of experts. Despite the significant progress in TDT diagnosis and treatment over the past decades that has dramatically improved patients' prognosis, its management remains challenging. On one hand, diagnostic and therapeutic advances are not equally applied to all patients across the world, particularly in several high-prevalence eastern regions. On the other, healthcare systems in low-prevalence western countries that have recently received large numbers of migrant thalassemia patients, were not ready to address patients’ special needs.

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the world by promoting education, research, awareness, and advocacy. One of the main actions of TIF is the development and dissemination of clinical practice guidelines for the management of thalassemia and other hemoglobinopathies. TIF guideline documents are drafted and reviewed by world experts in the field and are periodically updated to incorporate the continuous advances in the diagnosis and management of patients with hemoglobinopathies.

In 2021, the fourth edition of the guidelines for the management of TDT were published by TIF. The full version of the guidelines provides detailed information on the management of patients with TDT and the clinical presentation, pathophysiology, diagnostic approach, and treatment of disease complications or other clinical entities that may occur in these patients, while also covering relevant psychosocial issues. The present document is a summary of the 2021 TIF guidelines that focuses mainly on clinical practice issues and recommendations. Where appropriate, the level of evidence supporting certain recommendations is provided and classified as (A) data derived from multiple randomized clinical trials or meta-analyses; (B) data derived from a single-randomized clinical trial or large non-randomized studies; (C) expert consensus or opinion and small studies, retrospective studies, registries.

DIAGNOSIS OF THALASSEMIA

The term “thalassemia” refers to a group of blood diseases characterized by decreased or absent synthesis of one or more of the normal globin chains. According to the chain whose synthesis is impaired, the thalassemias are called α, β, γ, δβ, or εγδβ thalassemias. Most thalassemias are inherited as recessive traits. The most clinically relevant types are α and β thalassemias, resulting from the decrease of one of the two types of polypeptide chains (α or β) that form the normal adult human hemoglobin molecule (Hb A, α2β2). Based on their clinical severity and transfusion requirement, thalassemia syndromes can be classified phenotypically into transfusion-dependent thalassemias (TDTs) and nontransfusion-dependent thalassemias (NTDTs) as shown in Figure 1.

The diagnosis of thalassemias relies on measuring red blood cell indices and hemoglobin analysis and assessing the clinical severity of anemia. Molecular genetic testing may be useful for predicting the clinical phenotype and enabling presymptomatic diagnosis of at-risk family members and prenatal diagnosis. The diagnostic criteria for thalassemia and hemoglobinopathies are summarized in Figure 2.

Key points and recommendations

- Diagnosis of thalassemia should be considered in all those who have hypochromic microcytic anemia (A).
- In the diagnostic work-up for hypochromic microcytosis, iron deficiency anemia should always be excluded (A).
- Molecular analysis is not required to confirm the diagnosis of a β carrier, but it is necessary to confirm the α thalassemia carrier status (A).
- An α gene triplication or quadruplication should be taken into consideration in heterozygous β thalassemia subjects with a thalassemia intermedia phenotype (A).
- The hematological parameters including red-cell indices and morphology, followed by separation and measurement of Hb fractions are the basis for the identification of β thalassemia carriers (A).
- Detection of Hb H as Hb H inclusion body in a peripheral blood film using supravital staining (brilliant cresyl blue) is the hallmark of this condition (A).
- β globin sequence analysis may be considered first if the affected individual is not of an ancestry at high risk or if targeted analysis reveals only one or no pathogenic variant (B).
- α thalassemia are mainly due to deletions of different length and they can be detected with methods such reverse dot blot (RDP), Gap-polymerase chain reaction (PCR), or multiplex ligation-dependent probe amplification (MLPA) (B).
- Methods that may be used to detect rare or unknown deletions include quantitative PCR, long-range PCR and, above all, MLPA (B).

TREATMENT OF THALASSEMIA

Blood transfusions

The aim of blood transfusion in thalassemia is to deliver an effective and safe transfusion regimen while minimizing the burden of transfusion therapy on everyday life.

An effective transfusion regimen will result in (i) good growth and development; (ii) good energy levels, (iii) sufficient suppression of intra and extramedullary hematopoiesis. A safe transfusion regimen will (i) use a product that is collected, tested, selected, issued, and administered adherent to established quality and safety regulations and guidance; (ii) be administered by staff trained in blood transfusion; (iii) involve informed patient consent; (iv) be performed in a system with a good hemovigilance structure.
Blood transfusion therapy is decided upon the following criteria:

- Confirmed diagnosis of thalassemia.
- Laboratory criteria:
  - Hemoglobin level (Hb) <70 g/L on 2 occasions, > 2 weeks apart (excluding all other contributory causes such as infections) or
  - Clinical criteria irrespective of hemoglobin level:
    - Significant symptoms of anemia
    - Poor growth/failure to thrive
    - Complications from excessive intramedullary hemopoiesis such as pathological fractures and facial changes
    - Clinically significant extramedullary hemopoiesis

Key points and recommendations

- The diagnosis of thalassaemia should be confirmed with appropriate clinical and laboratory methods before the onset of transfusions (A).
- An informed consent should be obtained for blood transfusion therapy.
- Hemovigilance and adverse events reporting should be applied, as they key to the safety of blood transfusions.
- Careful donor selection and screening should be used, favoring voluntary, regular, nonremunerated blood donors (A).
- Before first transfusion, extended red-cell antigen typing of patients at least for D, C, c, E, e and Kell (A) and if available a full red-cell pheno/genotype should be performed.
- At each transfusion, ABO, Rh(D) compatible blood should be administered. Choosing units compatible for ABO, C, c, E, e, and Kell antigens is highly recommended (A).
- Before each transfusion, screening for new antibodies and an IAT cross-match should be performed, or in centers that meet regulatory requirements, an electronic cross-match should be performed where allowed (A).
- Leucodepleted packed red cells should be used where available. Prestorage filtration is strongly recommended, but blood bank pretransfusion filtration is acceptable. Bedside filtration is only acceptable if there is no capacity for prestorage filtration or blood bank pretransfusion filtration (A).
- Washed red cells should be used for patients who have severe allergic reactions (A).
- Red cells should be stored in CPD-A within 1 week of collection and in additive solutions for less than 2 weeks of collection where available (A).
- Transfusions should be performed every 2–4 weeks, maintaining pretransfusion hemoglobin above 90–105 g/L or up to 110–120 g/L for patients with cardiac complications (A).
- A record of red-cell antibodies, transfusion reactions and annual transfusion requirements should be kept for each patient (A).
- The post-transfusion hemoglobin should be kept below 140–150 g/L (A).

Iron overload and iron chelation

Iron overload occurs when iron intake is increased over a sustained period of time, either as a result of red blood cell transfusions or increased absorption of iron through the gastrointestinal (GI) tract. Both of these occur in thalassemias, with blood transfusion therapy being the major cause of iron overload in TDT and increased GI absorption being more important in NNTDT. When TDT patients receive regular blood transfusion, iron overload is inevitable because the human body lacks a mechanism to excrete excess iron. Iron accumulation is toxic to many tissues, causing heart failure, cirrhosis, liver cancer, growth retardation, and multiple endocrine abnormalities.
Diagnosis and monitoring of iron overload

The diagnosis and monitoring of iron overload are based on the complementary use of the following parameters:

**Serum ferritin**

Serum ferritin (SF) concentration is measured at least every 3 months (1–3 months). Target value is currently between 500 and 1000 μg/L. Measuring the trends in SF is a more reliable indicator for adjusting therapy than the use of single values.

**Liver iron concentration**

Liver iron concentration (LIC) is measured by magnetic resonance imaging (MRI)-based methods. LIC should be assessed using an externally validated and standardized MRI technique. Ideally, the same technique, if possible, should be used for each individual patient. Normal LIC values are up to 1.8 mg/g dry weight (wt). The risk of side effects is much lower even below 3–5 mg/g dry wt, while with levels of up to 7 mg/g dry wt are seen in some nonthalassaeic populations without apparent adverse effects. Sustained high LIC above 15 mg/g dry wt have been linked to worsening prognosis, liver fibrosis progression, or liver function abnormalities. LIC can also be used to calculate total body iron using the formula: total body iron (in mg/kg body wt) = 10.6 × LIC (in mg/g dry wt).

The frequency of LIC assessment should be guided by its level and its rate of change:

- Stable levels in the range 3–7 mg/g dry weight (dw): Every 1 or 2 years
- Levels >7 mg/g dw: yearly
- Levels falling rapidly or <3 mg/g dw: 6–12 monthly

**Myocardial iron**

Myocardial iron is assessed by T2* cardiac MRI, using an externally validated protocol and software, which should undergo periodic external calibration. The frequency of cardiac MRI scan should be guided by myocardial iron level, for example:

- Stable T2* >20 ms: 2 yearly
- T2* 10–19 ms: yearly
- T2* <10 ms: 6 monthly

It is particularly important to measure cardiac function when cardiac iron is high (eg, T2* <10 ms), as this is associated with a high risk of deteriorating function and heart failure and requires urgent intesification of chelation.

**Target organ function and other parameters**

In addition to measuring SF, LIC and cardiac T2*, target organ function, including cardiac, hepatic, and endocrine function, should monitored and patients should be regularly screened for iron-mediated damage including cardiac dysfunction, diabetes, hypothyroidism, hypoparathyroidism, and hypogonadal hypogonadism.

Additional parameters of iron overload include:

- 24 hours urinary iron estimation, not widely used in routine monitoring;
- plasma nontransferrin bound iron and labile plasma iron, the value of which as a guide to routine treatment or prognosis has not yet been established.

Monitoring of iron overload is essential in establishing effective iron chelation regimes, tailored to individuals’ specific needs. In addition to SF, LIC and myocardial T2*, to guide chelation therapy it is essential to:

- document the age of onset of regular transfusions and iron chelation therapy for each patient;
- maintain an annual record of blood usage (ml/kg packed red cells) and daily iron loading (mg/kg/day) for each patient.
Iron chelation therapy

The aims of iron chelation therapy are as follows:

- **Prevention therapy**: to maintain safe levels of body iron at all times, by balancing iron intake from blood transfusion with iron excretion by chelation (iron balance).
- **Rescue therapy**: to remove excess iron stored in the body.
- **Emergency therapy**: to urgently intensify iron chelation in case of iron-induced heart failure.
- **Dose adjustment of therapy**: to adjust dosing and treatment regimens to changing circumstances identified by careful monitoring of body iron and its distribution; monitoring is important to avoid: (a) under-chelation with increased iron toxicity; or (b) over-chelation and increased chelator toxicity.
- **Adherence to therapy**: to adhere to prescribed regular regimen; intermittent high-dose chelation can induce negative iron balance but does not provide continuous protection from labile iron and also risks increased toxicity from the iron chelator.

Three iron chelators are currently licensed for clinical use, deferoxamine, deferiprone, and deferasirox; their properties and indicated doses are reviewed in Table 1.

**Key points and recommendations**

- Uncontrolled transfusional iron overload increases the risks of heart failure, endocrine damage, liver cirrhosis, and hepatocellular carcinoma (B).
- Liver iron concentration can be used to calculate total body and serum ferritin is an approximate marker of LIC (B).
- MRI-based measurements should be performed by standardized and externally validated techniques, ideally in validated centers (C).
- Chelation therapy is an effective treatment modality in improving survival, decreasing the risk of heart failure, and decreasing morbidities from transfusion-induced iron overload (A).
- Chelation therapy at the correct doses and frequency can balance iron excretion with iron accumulation from transfusion (A).
- Absolute change in total body iron in response to chelation can be calculated from change in LIC (B).
- Direction of change in body iron in response to transfusion and chelation can usually but not always be estimated from the trend in serum ferritin (B).
- Prevention of iron accumulation using chelation therapy is preferable to rescue treatment because iron-mediated damage is often irreversible, and removal of storage iron by chelation is slow—particularly after it has escaped the liver (B).
- Response to chelation is dependent on the dose applied and the duration of exposure (A).
- Response to chelation is affected by the rate of blood transfusion (B).
- Cardiac iron accumulates later than liver iron, and is rare before the age of 8 years, affecting a subset of patients (B).
- Chelation of storage iron from the liver tends to be faster than from myocardium (B).
- Cardiac storage iron concentration is directly related to the risk of heart failure, which can be reliably estimated by MRI (eg, cardiac T2*), provided the center performing the measurement uses a validated method that has been independently calibrated (B).
- Chelation can reverse iron-mediated cardiac dysfunction rapidly (within weeks) by rapid chelation of labile iron, if 24 hours chelation cover is achieved (A).
- Chelation therapy removes myocardial storage iron slowly (months or years) (A).
- Over-chelation increases side effects from chelation therapy, and doses should therefore be decreased as serum ferritin or liver iron levels fall (demonstrated most clearly with DFO) (B).
- The optimal chelation regime must be tailored for the individual and will vary with their current clinical situation.
- Chelation therapy will not be effective if it is not taken regularly—a key aspect of chelation management is to work with patients to optimize adherence (B).

**Novel therapies**

An improved understanding of the pathophysiology of β-thalassemia has paved the way for the development of novel therapies. These can be classified into two major categories based on the underlying pathophysiology that they address:

- correction of the δβ globin chain imbalance by hematopoietic stem cell transplantation or gene therapy;
- targeting ineffective erythropoiesis or iron dysregulation by novel agents such as luspatercept.

**Hematopoietic stem cell transplantation**

Up to date, allogeneic hematopoietic stem cell transplantation (HSCT) is the only and curative treatment option for thalassemia major, with more than 3000 HSCTs performed worldwide.22–24 In a large EBMT survey of 1061 cases of MSD HSCT performed between 2000 and 2010 in 132 centers in 28 countries with a median patient age of 7 years, long-term overall survival (OS) and thalassemia-free survival (TFS) were 91% and 83%, respectively.25 In recent years, a number of factors including improved conditioning regimen, improved prevention of graft versus host disease (GvHD) and more effective antibacterial, antiviral, and antifungal treatment resulted in a significant improvement of outcomes for HSCT with a cure of thalassemia achieved in 80% to 90% of patients today.

**Key points and recommendations**

- HSCT should be offered to thalassemia patients and their parents at an early age, before complications due to iron overload have developed, if an HLA identical sibling is available.
- Either bone marrow or cord blood from an HLA identical sibling can be used.
- A matched unrelated donor can be used, provided that high compatibility criteria for both HLA class I and II loci are present.
- Haploidentical HSCT in thalassemia can be considered in experienced HSCT centers in the context of well-designed clinical trials.
- Myeloablative conditioning regimens should always be used for standard transplantation.
- Post-transplant care should include all transplant and thalassemia related complications.
- In thalassemia patients, HSCT is cost-effective when compared to life-long supportive therapy.

**Gene therapy**

For years, the only curative treatment has been allogeneic bone marrow transplantation, limited however by its availability only to young patients with a well-matched donor and the requirement for long-term immunosuppression to prevent or treat transplant-related immunological complications such as GvHD and rejection, carrying a nonnegligible risk of mortality.26 Gene therapy aspires to provide cure for thalassemia through the manipulation of the genome of hematopoietic stem cells, thus compensating for the inadequate or faulty function of mutated genes. This can be achieved by30,31:

- **gene addition** via a semirandom insertion of a healthy copy of the therapeutic gene into the cells using viral vectors or
- **gene editing** via a precisely directed mutation that repairs the gene in situ or induces a disease-modifying effect (ie, reactivation of Hb F synthesis) using site-specific nucleases.
Among the novel gene therapies, lentiviral vector gene therapy is the most mature intervention, shown to provide clinical efficacy and safety as a one-off, life-changing treatment. Nevertheless, the long-term safety and sustainability of the response needs also to be demonstrated; thus treated patients are followed in follow-up trials lasting overall 15 years. Precision medicine with genome editing tools has enabled overcoming of some of the obstacles associated with gene addition gene therapy; however, clinical experience with gene editing is currently limited and more clinical data and large-scale trials are needed to demonstrate that gene editing is a potential safe and curative treatment for hemoglobinopathies. The recent clinical data of patients with β genotypes create optimism for Zyglo's LentiGlobin vector approach.33,34

Key points and recommendations

Although waiting for the long-term clinical data on gene therapy for β-thalassemia, currently and on the basis of existing indications, patients with β-thalassemia major have potentially the following options for treatment:

- allogeneic hematopoietic stem cell (HSC) transplantation: young patients (≤17-year-old) with a β' or β genotype having an HLA-compatible sibling or a 10/10 matched volunteer donor.
- gene therapy with Zyglo: young patients in the 12- to 17-year-old age group with a β genotype who do not have an HLA-compatible sibling donor.
- gene therapy with Zyglo: patients in the 17- to 55-year-old age group with a β genotype who do not have severe comorbidities and are at-risk or ineligible to undergo an allo-HSC transplant but can otherwise undergo an autologous gene therapy procedure with an acceptable risk.

Targeting ineffective erythropoiesis—luspatcept

Luspatcept (ACE-536) is a recombinant fusion protein that binds to specific ligands of the TGF-β superfamily and enhances erythroid maturation. It is the most recently approved therapy [Food and Drug Administration (FDA) and European Medicines Agency (EMA)] for the management of TDT. The approval of luspatcept was based on the results of the phase 3 BELIEVE trial, which showed that subcutaneous luspatcept led to a reduction in transfusion burden of at least 33% from baseline during weeks 13 through 24 plus a reduction of at least 2 red-cell units over this 12-week interval in significant more patients than placebo (21.4% versus 4.5%); the percentage of patients with a reduction in transfusion burden of at least 33% during any 12-week interval was also greater with luspatcept (70.5% versus 29.5%). Parallel reductions in SF levels were further observed. Adverse events more common with luspatcept compared with placebo and included bone pain, arthralgia, dizziness, hypertension, and hyperuricemia. Hypertension developed in 10.7% and thromboembolic events in 3.6% of patients. Data on the long-term use of luspatcept, its real-life application and its use in the pediatric population are awaited. More details on this drug can be found in the full text of these guidelines.

Key points and recommendations

- Luspatcept can be considered for:
  - Patients who require regular red blood cell transfusions,
  - ≥18 years of age.
- The recommended starting dose of luspatcept is 1 mg/kg once every 3 weeks by subcutaneous injection.
- If the predose hemoglobin level is ≤115 g/L and is not influenced by recent transfusion, consider delaying dosing of luspatcept until the level is ≤110 g/L.
- Before administration of luspatcept, hemoglobin level, and liver function tests including alanine transference and aspartate transference levels should be monitored to ensure proper dosing and metabolism of the medication.
- If a TDT patient does not achieve a reduction in red-cell transfusion burden after at least 2 consecutive doses (6 weeks) at the 1 mg/kg starting dose, increase the luspatcept dose to 1.25 mg.
- If a patient experienced a response followed by a lack of or lost response to luspatcept, consider initiating a search for causative factors.
- Luspatcept should be discontinued if a patient does not experience a decrease in transfusion burden after 9 weeks of treatment (administration of 3 doses) at the maximum dose level or if unacceptable toxicity occurs at any time.
- It is important to monitor any TDT patient receiving luspatcept for signs and symptoms of thromboembolic events and initiate treatment accordingly.
- Blood pressure should be monitored before each administration of luspatcept.
- As no data are currently available on luspatcept use in pregnant women, all pregnant women should be advised of the potential risk to a fetus.
- Safety and efficacy of luspatcept in pediatric patients has not yet been established and its use in pediatric patients in therefore not currently recommended.

DIAGNOSIS AND MANAGEMENT OF THALASSEMA-RELATED COMPLICATIONS

Cardiovascular disease

Cardiovascular (CV) complications represent the leading cause of mortality in patients with thalassemia, including TDT and NTDT. In contemporary cohort studies, however, CV mortality has significantly declined, reflecting the overall mortality reduction observed in the same cohorts achieved by the effective implementation of modern diagnostic and therapeutic modalities, including magnetic resonance imaging (MRI)-guided chelation therapy with the use of the T2* technique. This progress is not, however, the case in thalassemia populations with limited access to modern therapy and therefore the global burden of CV disease in thalassemia remains high.

The pathophysiology and phenotypes of CV disease depend on the interaction between the main disease and the applied therapy. A wide range of CV abnormalities are seen in patients with thalassemia. The spectrum of cardiac disease includes left or right ventricular dysfunction, with or without heart failure, pulmonary hypertension, tachyarrhythmias such as atrial fibrilation, bradyarrhythmias such as atrioventricular block, valvular disease, pericarditis, and myocarditis. Further CV disorders include, thromboembolic events, resulting from either venous or arterial thrombosis, cerebrovascular disease, manifested as either ischemic or hemorrhagic stroke, and vascular abnormalities, including endothelial dysfunction and increased arterial stiffness. Among the above disorders, iron overload cardiomyopathy has long been the main form of heart disease in thalassemia patients treated with transfusions but is now being effectively prevented and managed with MRI-guided contemporary chelation therapy.

Regular CV assessment should be part of a multidisciplinary monitoring programme and should ideally be performed by or in consultation with clinics or physicians with experience in CV disease in hemoglobinopathies and in close collaboration with the attending thalassemia physician. Regular CV monitoring should be performed on an annual basis in all thalassemia patients, regardless of the presence of history or symptoms of CV disease. In the presence of CV disease, the frequency and content of CV assessment should be tailored to meet each
The prevention and treatment of CV disease in thalassemia consists of two pillars: the optimization of disease-specific therapy (blood transfusions, iron chelation, multidisciplinary care) and cardioactive therapy in the presence of CV disease (Figure 4). A lifestyle that promotes CV health is an important part of CV prevention, whereas the particularities of CV disease in thalassemia should be taken under consideration in the management of patients. Most importantly, cardiac dysfunction and heart failure may be reversible by timely therapy.

**Key points and recommendations**

- Regular CV assessment should be part of a patient’s multidisciplinary monitoring programme (C).
- CV assessment and management should ideally be performed by or in consultation with clinics or physicians with experience in CV disease in hemoglobinopathies and in close collaboration with the attending thalassemia physician (C).
- MRI T2* guided chelation therapy represents the best available approach to prevent cardiac dysfunction related to iron overload (B).
- In places lacking CMR T2* assessments, worsening of LV function in serial echocardiograms in transfusion-dependent patients may be used as a red flag for iron toxicity and should prompt aggressive and sustained escalation of chelation therapy (B).
- Echocardiographic screening for pulmonary hypertension should be part of annual CV assessment. Patients having a TRV greater than 3 m/s should undergo cardiac catheterization to confirm the diagnosis of pulmonary hypertension if a proximate cause cannot be identified and corrected (B).
- Intensified chelation therapy, preferably with two-drug combinations, is required for thalassemia major patients with cardiac iron overload, with or without overt cardiac dysfunction or heart failure (B).
- Diagnosis of CV disease should prompt optimization of disease-specific therapy in addition to cardioactive treatment (C).
- Screening and treatment of endocrine and metabolic comorbidities is crucial for the prevention and management of CV disease (C).
- Management of cardioactive therapies must account for a patient’s unique physiology compared with the general population (C).
- Cardiac abnormalities, including ventricular dysfunction, heart failure, and arrhythmias, are often reversible following intensification of disease-specific therapy, albeit after several weeks or months (C).
- Lifestyle choices that promote CV health (absence of smoking, physical exercise, weight control, healthy diet) should be vigorously promoted in thalassemia patients (C).

**Liver disease**

Iron overload and viral hepatitis are the two main causes of liver disease in patients with thalassemia leading to chronic inflammation, fibrosis and ultimately to cirrhosis. In iron overload states, unbound iron accumulates in the hepatocytes and leads to severe oxidative stress with overproduction of toxic ROS, which lead to severe hepatic inflammation and fibrosis. LIC estimated by MRI R2 or R2* is currently the method of choice to monitor iron overload specifically in this organ. Current iron chelators reduce significantly LIC and slow down liver fibrosis.

Hepatitis C virus (HCV) or hepatitis B virus (HBV) may further accelerate liver fibrosis in thalassaemic patients. Adequate blood donor checking, widespread vaccination for HBV and access to antiviral treatments are mandatory to minimize this risk. More specifically, a new generation of highly effective and safe direct-acting antiviral agents (DAA) have currently replaced interferon-based regimens in the HCV treatment landscape and have simplified treatment approaches for thalassaemic patients with chronic HCV infection.

The achievement of prolonged survival of thalassaemic patients in recent years is associated with effective iron overload treatments and lower risk of heart disease. However, there are several reports of a recent increase in hepatocellular carcinoma (HCC) incidence. Based on this observation, biannual screening for HCC with ultrasound for all thalassaemic patients is strongly recommended.

**Key points and recommendations**

- MRI R2 or R2* is the method of choice to assess liver iron concentration (LIC) and monitor chelation therapy effectiveness (A).
- Deferoxamine, deferasirox, and deferasirox are effective in decreasing total body iron burden as well as LIC (A).
Screening for HCV and HBV-chronic infection is recommended in thalassaemic patients (A).

Vaccination against hepatitis B is recommended in all patients with thalassemia who are seronegative for HBV markers (A).

If anti-HCV antibodies are detected, the presence of HCV-RNA in serum or plasma should be determined to identify patients with chronic infection (A).

The new IFN-free, ribavirin-free direct-acting antiviral drugs for hepatitis C are effective and safe in patients with thalassemia (B).

Serum transaminases, HBV DNA levels and liver fibrosis assessment by transient elastography are the main tools to guide treatment decision in HBV-chronic hepatitis (A).

Oral nucleoside and nucleotide analogues are well tolerated and effective drugs for HBV-chronic hepatitis, although loss of HbsAg remains a rare event (A).

Biannual ultrasound screening for hepatocellular carcinoma should be performed in all thalassaemic patients (B).

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Growth abnormalities

Growth failure in TDT has persisted despite major therapeutic advances. In the past, prevalence of growth failure and short stature in children with thalassemia varied from 30% to 60%. In the current era, the adherence to modern transfusion and iron chelation protocols and avoidance of iron chelator overdosage has clearly reduced the risk of short stature and may have potentially enhanced endocrine development in children with TDT.

The pathogenesis of growth failure is multifactorial. Key contributing factors to stunted growth in patients with TM include chronic anemia, transfusion-related iron overload, and chelation toxicity. Other contributing factors include hypothyroidism, hypogonadism, GH deficiency/insufficiency, zinc deficiency, chronic liver disease, under-nutrition, and psychosocial stress.

Patients’ growth should be assessed every 6 months by accurate measurement of standing and sitting height and weight. Diagnosis requires careful clinical evaluation to establish:

- Short stature—height below the third centile for sex and age (based on national growth charts), or
- Slow growth rates—growth velocity expressed in cm/year, below 1 SD for age and sex (based on growth velocity charts), or
- Signs of other pituitary hormone deficiencies (eg, gonadotrophins, growth hormone, central hypothyroidism)
- Signs of other possible causes of retarded growth (nutritional deficiencies, chronic hepatic disease, chronic heart failure).

Management consists of optimizing blood transfusion; improving nutrition with high caloric balanced diet. Optimizing iron chelation can be achieved by using the new oral chelators or intensive and combined chelation therapy in patients with severe iron overload. Also, an early diagnosis and treatment of associated endocrinopathies are important. Recombinant GH (rhGH) treatment is not always as effective as expected in non-thalassaemic children with GHD.
velocity attained after exogenous GH administration in children with thalassemia is reported to be lower than that seen in children with primary GH deficiency, possibly due to GH insensitivity (B).

- Essentially there is no pathognomonic clinical feature to lead to suspicion of GH deficiency in adults (B).

**Endocrine complications**

Endocrine abnormalities are the most common complications of TDT. Prevalence varies because of the different levels of treatment followed by centers across the world, particularly the severity of the defective genetic background, the levels of hemoglobin and the level of iron load in various patient groups. Another contributing factor is that of the increased survival to adulthood.51,53

Endocrine-related complications include the following:

- **Delayed puberty and hypogonadism:** Hypogonadotropic hypogonadism ranges from 50% to 100%. Adult-onset hypogonadism (AOH) in TDT patients ranges between 8.3% and 12%. These complications result mainly from iron overload (A).

- **Hypothyroidism:** It varies from 6% to 35%.

- **Impaired glucose tolerance and insulin-dependent diabetes mellitus (IDDM):** It increases with age and varies from 10% to 24%. The etiology of IDDM is multifactorial (genetic factors, insulin deficiency, insulin resistance, and liver dysfunction secondary to viral hepatitis).

- **Hyperparathyroidism:** It varies from 1% to 19%. Most patients show a mild form of the disease.

- **Adrenal insufficiency:** "Biochemical adrenal insufficiency" varies up to 45%, but clinical adrenal insufficiency is rare.

**Key points and recommendations**

- Endocrine complications are very common in multitransfused TM patients (A) and periodic evaluation for endocrine complications should be carried out in TDT patients with iron overload, particularly after the age of 11 years (B).

- Sub-clinical hypothyroidism (basal TSH 5–8 mUI/ml) requires regular follow-up and optimizing chelation therapy (B).

- Normalization of total body iron load with intensive combined chelation (desferrioxamine plus deferasirox) can reverse endocrine complications of TDT (B).

- Monitoring of growth, pubertal development, reproductive ability, and endocrine functions in general are essential to achieve a good quality of life in TDT (B).

**Infectious disease**

Infections and their complications were previously the second commonest cause of death in TDT,26 but are currently becoming the leading cause of mortality in western countries due, in part, to a significant reduction in the deaths from iron-induced cardiac disease.24 There is a lack of properly controlled studies evaluating infections in thalassemia. The knowledge of infections depends more on anecdotal reports and experimental studies. The mechanisms of susceptibility to infections in thalassemia have yet to be clarified. The better understanding of underlying mechanisms and their impact on evolving infections, regional and community-based differences in infectious risks and preventative measures may contribute to a reduction in infection-related mortality in thalassemia.

**Key points and recommendations**

- Physicians must be aware of the potential life-threatening infections in thalassemia and patients should be educated to seek early care when fever develops.

- Control of iron homeostasis may have therapeutic benefit against infections.

- Temporary discontinuation of deferoxamine and prompt initiation of antibiotics are strongly advised; whereas, iron-loaded patients can continue to use synthetic oral iron chelators such as deferasirox during febrile episodes.

- Transfusion of prestorage leucodepleted red cells that have been stored <14 days may have therapeutic benefit against infections.

- Quality assurance guidelines and strict regulatory standards should be established for enhancing transfusion safety.

- Splenectomy indications and preventive measures for postsplenectomy risk of sepsis should be revisited.

**Bone disease**

Bone abnormalities are often seen in thalassemia patients and have become a major cause of skeletal complications. These skeletal unusual changes include decreased bone mineral density (BMD), spontaneous fractures, spinal deformities with compression of the vertebrae and nerves often causing severe pain and discomfort. Males with TDT seem to be more frequently and severely affected from bone disease, suggesting a possible gender variation.

Osteoporosis is a prominent cause of morbidity in patients with thalassemia major; it is present in approximately 40%–50% of patients.57 The pathogenesis includes genetic factors as well as endocrine complications (mainly hypogonadism), iron overload, bone marrow expansion, vitamin deficiencies, and lack of physical activity.58 These factors can lead to bone destruction through the increase of osteoclast function and/ or the reduction of osteoblast activity.

Management of thalassemia-associated osteoporosis consists of adequate calcium and vitamin D intake, sufficient iron chelation, hormone replacement, and inhibition of the osteoclast function mainly by bisphosphonates.56–58 Intravenous administration of pamidronate or zoledronic acid seems to be more efficient than oral bisphosphonates. Other novel agents such as the novel osteoclast inhibitor, denosumab, teriparatide, and the activin antagonist, sotatercept, are under investigation but their effects in TM-induced osteoporosis remains to be proven.56,59

**Key points and recommendations**

- Annual checking of BMD starting in adolescence is considered indispensable (C).

- Biochemical markers of bone metabolism that can be done every year: N-telopeptide of type I collagen (NTX), C-telopeptide of type I collagen (CTX), bone-specific alkaline phosphatase (bALP) (C).

- Physical activity must always be encouraged (C).

- Smoking should be discouraged (C).

- Adequate calcium intake during skeleton development can increase bone mass in adult life and in combination with administration of low doses of vitamin D may prevent bone loss and fractures (C).

- Early diagnosis and treatment of diabetes mellitus (C).

- Adequate iron chelation may prevent iron toxicity in the bone and sufficient blood transfusions may inhibit uncontrolled bone marrow expansion (A).

- Hormonal replacement where it is needed (A).

- Bisphosphonates should be given concomitantly with calcium and vitamin D and not for longer than 2 years (A).

**Splenomegaly and splenectomy**

Increased destruction of red blood cells by the reticuloendothelial system, in particular the spleen together with extramedullary
hemopoiesis result in splenic enlargement (splenomegaly). The main therapeutic rationale for splenectomy in transfusion-dependent patients with TDT has been to decrease blood consumption and transfusion requirements with the ultimate goal of reducing iron overload. However, current transfusion guidelines setting more adequate pretransfusional hemoglobin levels (90–105 g/L) have considerably reduced the incidence of splenomegaly and the need for splenectomy in TDT patients. Splenectomy should be restricted to certain indications in view of the increased risk of venous thrombosis and pulmonary hypertension, alongside overwhelming infections after splenectomy (Table 2). Splenectomy should be avoided in children less than 5 years of age because of a considerably greater risk of fulminant postsplenectomy sepsis.

Key points and recommendations

- The size of the spleen should be carefully monitored and recorded on physical examination and, as needed, by ultrasonography.
- Splenectomy is the recommended intervention to reduce excessive blood consumption and consequent severe iron overload. However, physicians should keep a guarded approach towards splenectomy because of the high disease burden associated with splenectomy. Current optimal transfusion regimens and iron chelation have considerably reduced the incidence of splenomegaly and iron overload in transfusion-dependent thalassemia patients.
- At the current time, we do not recommend splenectomy as a standard procedure in patients with thalassemia (C). There is large amount of evidence that links splenectomy to a variety of complications such as pulmonary hypertension, silent brain infarcts, venous thrombosis, and sepsis to name a few. Splenectomy should be considered in patients with thalassemia in three clinical scenarios: increased blood requirement that prevents adequate control with iron chelation therapy, hypersplenism, and symptomatic splenomegaly (C).
- When performing splenectomy, the laparoscopic approach seems to be the most favorable (B).
- The most frequent pathogens that cause infections in splenectomised patients are *Streptococcus pneumoniae*, *Haemophilus influenzae* type B, and *Neisseria meningitidis*; therefore, immune prophylaxis is recommended against these agents at least 2 weeks before the operation and repeated postoperatively as recommended. Additionally, an annual influenza vaccination is encouraged.
- Chemoprophylaxis with oral penicillin depends on the age of the individual and the treating physician’s opinion (C). Life-long prophylactic antibiotics with oral penicillins or macrolides should be offered to patients considered at constant high risk of pneumococcal infection, including those who have previously experienced an episode of sepsis or those suffering from concomitant hematological diseases or other immune system impairment.
- In current practice, due to strict transfusion and chelation protocols, the disease is very well controlled and we are seeing fewer splenectomies than before. Nevertheless, a large percentage of the thalassemia population (particularly in the older age group), is already splenectomised. These patients are at an increased risk of many disease-related morbidities and should be monitored closely.
- Adhere to current transfusion guidelines to prevent or delay splenomegaly.
- If splenectomy cannot be avoided, make sure immunization protocols are followed.
- Be aware of the dangers of thrombosis in the peri-and post-operative period.
- Discuss postsplenectomy chemoprophylaxis with patient/family and make sure they are aware of the dangers of postsplenectomy sepsis.
- Start appropriate parenteral antibiotics in case of suspected sepsis while awaiting culture results.

### Table 2. Indications for splenectomy in TDT

| INDICATION | COMMENT |
|------------|---------|
| Increased blood requirements that prevent adequate control with iron chelation therapy, having made sure that the increased requirements are not due to allo/auto-antibodies or blood loss. | Annual transfusion volume used to flag an increased blood requirement (200-275 ml/kg/yr red cells)*. However, if effective chelation therapy continues to be maintained despite increased transfusion, splenectomy may not be necessary. |
| Hypersplenism | Cytopenias causing clinical problems |
| Symptomatic splenomegaly | • Accompanied by symptoms such as left upper quadrant pain or early satiety. • Massive splenomegaly causing concern about possible splenic rupture. |

*Guidelines vary slightly as to what constitutes increased requirements: Physicians should make a decision based on all clinical parameters.

TDT = transfusion-dependent thalassemia.
Dental care

Thalassemia patients show a tendency for higher caries, plaque rates, gingivitis, and periodontitis scores than control subjects.67 Those may be attributed to poor oral hygiene, improper dietary habits, lack of dental awareness, reduced salivary flow rate, and neglected dental care. As patients with TDT develop certain disease-related orofacial changes, awareness of these changes is important for the proper dental care.68

Key points and recommendations

• Dentists need to be aware of the orofacial manifestations of thalassemia so that they can be identified early and appropriately managed.
• Patients with TDT are at increased risk of developing dental caries and periodontal disease.
• Patients should be maintained closely on a preventive programme with regular follow-up. Oral hygiene instructions, dietary advice, and preventive measures including prophylaxis, fluoride application, and fissure sealants should be implemented to minimize the need for invasive dental procedures.
• Close liaison with the hematology team is required to determine the potential complications when delivering invasive dental treatment and measures put in place to reduce risk.
• Predisposing factors for infections in thalassaemic patients include severe anaemia, iron overload, splenectomy, and a range of immune abnormalities. Guidelines regarding anti-biotic prophylaxis vary from country to country with some recommending prophylaxis similar to that used for the prevention of bacterial endocarditis.
• Correction of drifted maxillary anterior teeth and increased overjet should be undertaken to improve aesthetics, reduce susceptibility to trauma, avoid gingival inflammation, and improve functional ability. It is recommended that orthodontic treatment be initiated as early as possible, concentrating on preventive and interceptive approaches.
• All patients should ideally have a comprehensive dental assessment with their local dentist prior to the commencement of bisphosphonate therapy, to ensure that they are as dentally fit as feasible and be aware of bisphosphonate-related complication (bisphosphonate-related osteonecrosis of the jaw). Preventive dental advice should be given, emphasizing the importance of reporting any symptoms such as loose teeth, pain, or swelling, as soon as possible.

Lifestyle and quality of life

Although TDT is burdensome conditions, requiring life-long treatment and close follow-up and can often be complicated by complications arising from many different systems, patients with optimally treated thalassemia can now enjoy a near-normal life and lifestyle, and experience full physical and emotional development from childhood to adulthood.

Key points and recommendations

• A holistic approach to patient care includes recognition of the need for social integration and a “normal” life.
• The treating physician should be able to provide advice on lifestyle issues.
• Physical activity should be encouraged but the condition of each individual patient should be recognized and comorbidities identified. Cardiovascular assessment and ergometry if available should be undertaken before recommending exercise and activity.
• Clinic and transfusion times should be flexible and take into consideration the patient’s commitments, such as their education and work. Availability of some evening and weekend clinic and transfusion sessions is highly recommended.
• Liaison with teachers and employers to provide understanding of the condition and its management may be necessary.
• Routine monitoring of growth is necessary, and nutritional factors such as caloric intake and micronutrient deficiencies should be considered in instances of poor growth. The services of a dietician are helpful in this respect.
• Zinc supplements may be given in cases of deficiency, poor growth and reduced bone mass, but are not recommended as routine for all patients.
• Dietary iron restriction is recommended in patients on low transfusion regimens with low pretransfusion hemoglobin.
• Calcium and vitamin D supplements are recommended for all patients at a dose of 2000 IU per day, along with measurements of vitamin D levels every 6 months. A diet high in calcium is also recommended (through milk, fish, cheese, etc.).
• Folic acid supplements of up to 1 mg/day are needed for all patients with low hemoglobin levels. Supplementation for all patients may be considered, since the risk of thrombosis may be reduced and toxicity is low.
• A diet rich in foods with high vitamin E content, such as eggs and vegetable oils is recommended. Prolonged use of supplements requires further research.
• Vitamin C supplements are recommended, in conjunction with deferoxamine infusions at a dose of 2–3 mg/kg/day, or if deficiency is proven.
• L-carnitine may be beneficial at a dose of 50 mg/kg/day, although caution should be exercised in patients with thyroid dysfunction.
There is insufficient evidence on any long-term benefits from wheat grass.

- Silymarin at a dose of 140 mg three times daily is recommended if liver involvement is detected, and on consultation with a hepatologist.

- Consumption of alcohol and tobacco and substance abuse should be avoided.
- A quality of life assessment should form part of the regular evaluation of each patient's progress, and may usefully also highlight issues in service delivery.
### Table 4.
General Timetable for Clinical and Laboratory Investigation.

| INITIAL ASSESSMENT / DIAGNOSIS | Routine Tests | Clinical Testing | Laboratory Testing | Imaging Testing |
|-------------------------------|---------------|------------------|---------------------|-----------------|
| **Laboratory Tests**<br>Recommended before 1st Blood Transfusion | **Children and Adolescents** | **Adults** | **As Clinically Indicated** |
| CBC Hb, Indices; Nucleated Red Cells; Platelets | Monthly 3-Monthly 6-Monthly 12-Monthly | Monthly | | |
| | Height | Growth Velocity | Bone Deformity | | |
| | Weight | | Bone Age (for the first 10 years of life) | | |
| Molecular Mapping | | | | | |
| α-Globin Gene Cluster | Transfusion Reactions | Serum Ferritin | Zinc Levels | Holter ECG |
| β-Globin Gene Cluster | Weight | Transferrin Saturation | Magnesium Levels | Right Cardiac Catheterisation |
| ABO System | Blood Pressure | Transaminases | Vitamin Levels | (if Elevated TR)
| Rh System and Subgroups | Complete Blood Count | γ-GT, ALP, LDH | Cholesterol / Triglycerides | Liver Biopsy (Optional)
| HLA Type | Nucleated Red Cell Count | Bilirubin (t,d) | | Duplex Abdominal Ultrasound |
| G6PD Quantification | New Red Cell Antibodies | Total Protein | | MRI of Hypothalamic–Pituitary Region |
| **Clinical Tests**<br>(Repeat at each visit and/or as indicated) | Temperature | Albumin | | Ergospirometry |
| | Pulse | Blood Urea | | Echocardiography |
| | O2 Saturation | Creatinin | | MRI Pancreas |
| | | Uric Acid | | MRI Cardiac T2* |
| | | Electrolytes | | Thyroid Ultrasound |
| | | Calcium Ionised | | MRI Liver Iron (LIC) |
| | | Phosphate | | PCT |
| | | Fasting Sugar | | Abdominal Ultrasound |
| | | Creatine Kinase | | (screen for liver disease)
| | | | | DEXA Scan |
| | | | | Fibroscan (Liver Disease) |
| | | | | HAV Serology (only nucleated) |
| | | | | HBV Serology/PCR (only nucleated) |
| | | | | HCV Serology/PCR (only nucleated) |
| | | | | HIV Serology |
| | | | | OGT |
| | | | | Thyroid Function (>9yrs)
| | | | | Parathormone (>16yrs)
| | | | | LH-ICMA (>12-14yrs) |
| | | | | FSH (12-14yrs)
| | | | | Estradiol/Testosterone (10-14yrs) |
| | | | | IGF-1, IGF BP-3 |
| | | | | Cortisol | Test for Bone Formation |
| | | | | (after ACTH Stimulation) | (e.g. Bone Specific AP) |

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Psychological support

Overall, despite a general lack of large-scale, randomized, controlled trial evidence conducted with patients with thalassemia, there are innumerable cohorts of case controlled analytical studies to suggest that psychological well-being has an impact on adherence to treatment for chronic disease in general (B).72–74 In thalassemia, the published reports to demonstrate this linkage are mainly descriptive studies (C). A meta-analysis would suggest that more recent efforts are more towards B grade investigations (usually ancillary studies attached to robust controlled trials in other clinical areas). However, the lack of uniform instruments and standardized measurements weakens this assessment. The findings to date suggest that:

- Psychological well-being impacts on adherence to chelation treatment in TDT and hence on survival (C).
- Patients with thalassemia are vulnerable to experiencing psychological challenges (C).
- Patient-reported health outcomes show that oral chelation therapy has a beneficial impact, relative to parenteral chelation (B).
- Neuropsychological investigation of cognitive deficits shows that there are clear intellectual and psychopathological problems in a very limited number of thalassemia patients (B).
- Benefits of psychological support have been suggested using a variety of approaches (C) which include:
  - Targeting changes in institutional organization practices
  - Patient group sessions
  - Family therapy
  - Patient chelation camps
- In all chronic illness, continuity of comprehensive care across the lifespan is essential for long-term, beneficial health outcome (A). Institutional organizational support for multidisciplinary teams is essential (A). There is a growing body of evidence that highlight the problems associated with transition from pediatric care to adult internal medicine in inherited chronic disease (B). Rare and neglected diseases complicate resource allocation models and lead to notable health disparities (A). In thalassemia, these problems are known and reports from expert committees recommend addressing them, but there are no formal studies of the problems, much less any standardized evidence.

While A and B grade evidence for psychological support in thalassemia is scarce, experience in several large thalassemia centers strongly suggests that psychological well-being is key to adherence and to outcome.

- Expert psychological support has to be available at all centers specializing in thalassemia care (C).
- Additional psychosocial support provided by trained specialists (eg, social workers or family or child health specialists) should be tailored to the patient’s age
  - Children (in general, A, thalassemia C)
  - Adolescents – transition (in general, B, thalassemia C)
  - Older adults – pain issues (in general, A, thalassemia C)

Funding for clinical psychological support services could be more widely achieved if well-designed, multicenter, intervention studies using common standardized instruments were undertaken to evaluate the benefit of psychological and psychosocial support to treatment adherence. The use of established behavioral and social science approaches in such studies need to identify the active components of “psychological support” that are most applicable to patients with thalassemia.

Multidisciplinary care

The life-long and multigenerational nature of TDT imposes a multidisciplinary approach to patient’s care. As patients advance in years, the basic needs in blood transfusion and iron chelation, even if and when provided appropriately and in accordance to international standards, gradually become inadequate to sustain life, maintain well-being, and achieve social integration. For this reason, specialists in several medical disciplines including but not confined to heart, liver, and endocrine, are called upon to contribute by monitoring and offering proactive management of iron toxicity and organ dysfunction in their field of expertise.75 An example of the structure of interdisciplinary team for the care of hemoglobin disorders as extracted from some well-established European Reference Centres with successful patient outcomes is presented in the Table 3.60 A suggested timetable for the multidisciplinary clinical and laboratory investigation and follow-up of patients with TDT is provided in Table 4.

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