Review of endocrine complications in adult patients with β-thalassaemia major

Ploutarchos Tzoulis

UCL Medical School, Royal Free Hospital, London, UK

Abstract

Endocrine abnormalities are amongst the most common complications of β-thalassaemia major (TM). This is an overview of endocrinopathies of adult patients with β-thalassaemia major, excluding osteoporosis and fertility issues. This review will focus on emerging evidence in the last 5 years with regards to endocrinopathies in patients with TM.

Prevalence

In light of the significant increase of life expectancy of patients with TM in recent years, endocrinopathies have become even more topical. The prevalence of various endocrine disorders varies significantly between different studies. Numerous studies suggest that more than half of patients with TM, even in paediatric cohorts, have at least 1 endocrinopathy.1,6 The most recent study by Ang et al., including 92 adult patients with TM (median age 36 years) from the largest UK thalassaemia unit, showed much higher prevalence rates of endocrine disorders than previous studies. In this cohort, 75% of patients had at least one endocrinopathy, 67% had hypogonadism, 41% diabetes mellitus (DM), 17% hypoparathyroidism and 14% hypothyroidism.1 This difference was attributed to the much older age of this patient cohort in comparison to previous studies.1 This study, in keeping with other studies,2 showed also that increasing age is an independent risk factor associated with DM and hypogonadism. A past study, conducted in the same major thalassaemia unit in the UK in 1997, included 97 patients with mean age of 24 years and reported that 66% of patients had hypogonadism, 20% DM, 13% hypoparathyroidism and 10% hypothyroidism.3 It also demonstrated a clear association between genotype in TM major and hypogonadism as well as DM.2 Recent data of the French National Registry for β-TM showed that 55.2% of patients older than 24 years had hypogonadism, 11.2% diabetes and 18.0% hypothyroidism.2 The French Registry also found that the frequency of endocrine organ damage related to iron overload increased with age, with the occurrence of complications being rare in childhood.2 The largest study of prevalence of endocrine complications, including 3,817 patients with TM in 29 countries, showed lower prevalence rates than the recent UK and French data such as 40.5% for hypogonadism, 9.9% for glycaemic abnormalities, 6.9% for hypoparathyroidism and 3.2% for hypothyroidism.2 However, this large multi-centre study included also a large proportion (36%) of patients under the age of 16.

Longitudinal studies have reported that the prevalence of endocrine complications has declined in the last few decades thanks to more effective iron chelation and earlier age of first exposure to chelation treatment.8 However, recent studies1,2 have indicated increase in the prevalence rates of endocrinopathies as the mean age of cohorts and life expectancy have risen significantly. Overall, the prevalence of endocrine disorders in TM patients is affected by various factors such as age, degree and type of chelation, compliance with chelation, age of first exposure to chelation therapy, age of first transfusion, haemoglobin level attained before blood transfusion and genotype of TM.

Predictors / pathophysiology of endocrinopathies

Iron toxicity has been regarded as the most likely cause of development of endocrine disorders in patients with TM. A combination of pathologies, including direct tissue damage from iron deposition, damage from chronic anaemia, chelation of other essential elements such as zinc, direct toxicity of non-transferrin-bound iron through reactive oxygen species formation contribute to the development of endocrinopathies in this context.

A recent retrospective study1 on the associations between common endocrinopathies and iron load parameters (longitudinal data over a mean of 8 years) demonstrated that:

- There was a strong association between a history of myocardial T2 < 20 msec and hypogonadism as well as DM in keeping with similar findings in a cross-sectional study by Au et al. which also showed strong association of low myocardial T2 with hypoparathyroidism and hypothyroidism.3 Patients with hypogonadism and DM had median myocardial T2 values of 12.6 msec and 12.3 msec compared with 23.8 msec and 24.8 msec respectively in patients without those complications. Myocardial T2 < 20 msec was strongly associated with the occurrence of DM [OR 19.3; 95% CI 4.3 - 86.7; P < 0.001] and hypogonadism [OR 3.9; 95% CI 1.4 – 10.5; P=0.008].1
Increasing age was associated with DM and hypogonadism. Each 1 year increase in age was associated with 1.1 increased odds of developing DM [95% CI 1.0 – 1.2; P=0.002] and hypogonadism [95% CI 1.0 – 1.2; P=0.016].

Average 10-year serum ferritin levels > 1250 mcg/l and > 2000 mcg/l were significantly associated with DM [OR 14.8; 95% CI 2.4 – 90.0; P=0.003] and hypogonadism [OR 2.9; 95% CI 1.0 – 8.3; P=0.047] respectively. Some studies, but not all, have identified serum ferritin as a prognostic marker for progression to endocrine dysfunction.10,11

Liver T2 values as well as LIC (liver iron concentration) by MRI-R2 (Ferriscan) values were not associated with the development of endocrinopathies.

Therefore, myocardial iron loading is a good surrogate marker for significant iron overload in the pancreas and pituitary gland, as also shown in other studies which suggested that iron loading in the pituitary and pancreas precedes myocardial iron loading.12-14 Myocardial T2 < 20 msec is a good predictor of significant and prolonged iron overload in the pituitary and pancreas. Abnormal myocardial T2 should prompt intensification of iron chelation therapy and investigation and close monitoring for endocrinopathies.

**Reversibility of endocrine complications**

Endocrine complications of TM were in the past regarded, in general, irreversible. This concept has been recently challenged since there is high quality evidence from a randomised trial by Tanner et al. that combined therapy with desferrioxamine (DFO) and deferiprone (DFP) compared to desferrioxamine monotherapy can improve cardiac iron loading and cardiac function (an increase by 3% in ejection fraction over one year).15

A study by Farmaki et al., including 52 patients with a 5-7 year follow-up, demonstrated the efficacy of combined chelation therapy with DFO and DFP in reducing total body iron load and reversing as well as preventing endocrine complications. Patients on combination therapy had a statistically significant reduction of the total body iron load in comparison to previous DFO monotherapy, as indicated by ferritin, cardiac and liver iron. The proportion of patients with abnormalities in glucose handling such as DM, impaired glucose tolerance (IGT) and impaired fasting glycaemia (IFG) was reduced from 78% to 34%.16 Amongst 18 patients on thyroxine, 10/18 managed to discontinue thyroxine and 4/18 reduced the dose. Thus, intensive chelation therapy reversed most cases of compensated or subclinical hypothyroidism and also prevented any new cases of hypothyroidism. In addition, 50% of male hypogonadal patients achieved normal testosterone levels and were able to discontinue testosterone replacement therapy. Therefore, patients receiving intensive combined chelation reversed and prevented multiple endocrine complications through decrease of iron load which had accumulated over an extensive period.16

A different study by the same group17 assessed the effect of enhanced iron chelation treatment on glucose metabolism in patients with b-TM major. 41 patients, previously treated with DFO only, were switched to combined treatment with DFO and DFP. At baseline, 6/41 had DM, 15/41 IGT, 6/41 IFG and 14/41 had normal glucose handling. Combination therapy markedly reduced ferritin levels from 2991 ± 2093 mcg/l to 638 ± 1345 mcg/l (P < 0.001). Glucose responses were improved at all times during an oral glucose tolerance test (OGTT), particularly in patients in early stages of glucose intolerance. Out of 21 patients having at baseline IGT or IFG, more than half (12/21) restored normal glucose handling. Insulin secretion increased markedly in the overall group of patients, mainly due to the increases in insulin secretion in the IGT patients. Insulin sensitivity declined in all groups of patients, but this difference did not reach statistical significance. In total, the most significant improvement in glucose disturbances were found in patients with IGT and IFG.17

Another study, examining the effect of intensive chelation on endocrinopathies, found that intensive chelation therapy reversed hypothyroidism, while it did not have positive effect on glycaemic status or gonadal function. After a 4-year observational period, patients who developed new endocrine/metabolic complications were switched from DFO monotherapy to a combined intensive iron chelation regimen of DFO + DFP. After 24 months of intensive chelation with subsequent reduction of median ferritin from 1500 mcg/l to 569 mcg/l (P < 0.001), 8/12 patients on thyroxine were able to discontinue it and 4/12 reduced their thyroxine dose.11

A multi-centre retrospective cohort study, published in 2014, examined the incidence and progression of endocrine disorders during long-term treatment with deferasirox (Exjade) in a real clinical practice setting. This cohort included 86 patients with TM treated with once daily Exjade for a median duration of 6.5 years. The incidence of new endocrine complications was only 7% and included 5 new cases of hypogonadism and 1 new case of hypoparathyroidism. No new cases of diabetes or hypothyroidism were recorded. These findings suggested a low rate of new endocrine disorders and stabilisation of pre-existing disorders such as diabetes and hypothyroidism.18

**Glucose metabolism**

The prevalence of diabetes mellitus (DM) in patients with β-TM has been estimated to be between 3.2 and 41%.12,17,19 Abnormalities in glucose handling are relatively common complications in patients with TM. According to most recent diagnostic criteria published by American Diabetes Association in 2010, different states in glucose metabolism based on 75-gr oral glucose tolerance test (OGTT) are classified into:

- Diabetes (DM): Fasting plasma glucose (FPG) ≥ 7.0 mmol/l (≥ 126 mg/dl) or 2-hour plasma glucose ≥ 11.1 mmol/l (≥ 200 mg/dl)
- Impaired Glucose Tolerance (IGT): 2-hour plasma glucose 7.8 – 11.0 mmol/l (140 - 199 mg/dl)
- Impaired Fasting Glycaemia (IFG): FPG 5.6 – 6.9 mmol/l (100 - 125 mg/dl)
- Normal glucose metabolism: FPG < 5.6 mmol/l (< 100 mg/dl) and 2-hour plasma glucose < 7.8 mmol/l (< 140 mg/dl)

The pathogenesis of glycaemic abnormalities in β-TM is complex and multifactorial. It has been predominantly attributed to a combination of reduced insulin secretory capacity and insulin resistance. These patients are a very heterogeneous group with some individuals exhibiting mainly insulin deficiency and others predominantly insulin resistance. The traditional concept has been that the initial insult is insulin resistance compensated by hyperinsulinaemia. Then, pancreatic damage and insulin deficiency subsequently develops leading to DM. However, this is not always the sequence of events leading to development of DM. It has been shown that a defect in β-cell insulin secretion can be present early before the development of glucose intolerance, resulting from toxic effects of iron deposition in the pancreas.20 Overall, the interplay between liver siderosis, causing insulin resistance and pancreatic β-cell siderosis, causing insulin deficiency, leads to the development of diabetes. Also other factors play an important role such as hepatitis C viral infection, autoimmunity, family history of diabetes mellitus and genetic factors.21

According to the UK standards for the clinical care of children and adults with thalassaemia published in 2008, all patients should be screened with OGTT annually from puberty or from the age of 10 years if there is a family history of DM. Annual screening becomes even more
important in the light of evidence showing that intensive combined chelation regimen in the early stages of glucose abnormalities can improve insulin secretion and normalise glucose metabolism.17

Diagnosis of IFG or IGT indicates a pre-diabetic state which, if not managed appropriately, will progress to diabetes. Management of these patients should include intensive chelation therapy as well as lifestyle modification. Lifestyle modification in the form of regular exercise (for example walking 150 minutes/week) and weight loss of 5-10% of body weight has been shown to reduce drastically the progression of pre-diabetes to diabetes in the general population.

Management of DM should be individualised. The 1st line treatment in most patients should be, besides lifestyle modification, oral antidiabetic agents. There is very limited published data on the efficacy and safety of these agents in patients with TM. The only drugs used in small studies in this context with good effect are metformin,22 glibenclamide23,24 and acarbose.25 Modern hypoglycaemic agents such as glitazones (insulin-sensitising agents), DPP-IV inhibitors (increasing intrinsic GLP-1 levels), injectable GLP-1 agonists (mediating glucose-dependent insulin secretion) and SGLT-2 inhibitors (increasing renal excretion of glucose) have not been studied in patients with TM, but are increasingly becoming the mainstay of treatment of DM in the general population because of low rate of hypoglycaemias and weight loss. Some patients will need insulin therapy.

Their glycaemic control should be assessed by periodical fructosamine estimation (with fructosamine target < 322 umol/l equivalent to HbA1c < 7.0%). They should perform home capillary blood glucose monitoring with a frequency based on their treatment.

All patients with DM should be monitored regularly for the development of complications. Their monitoring, on at least annual basis, should include:

- Measurement of urine albumin creatinine ratio
- Assessment of renal function
- Retinal screening
- Foot examination
- Measurement of blood pressure (BP target < 130/80 mmHg)
- Lipid profile with targets for total cholesterol < 4.0 mmol/l (< 155 mg/dl) and LDL < 2.0 mmol/l (< 77 mg/dl)

It is worth emphasising that patients with TM and diabetes can develop diabetes-related complications. Microvascular complications include diabetic nephropathy, retinopathy and neuropathy. Limited data exist about the frequency of these complications, for example the rate of diabetic retinopathy has been reported 13.6 – 26.0%.21,26 A prevalence of microalbuminuria of 13.2 – 55.0%21,27 has been recorded. With regards to macrovascular complications of diabetes, they include ischaemic heart disease, cerebrovascular disease and peripheral vascular disease. Recent study by Pepe et al. showed that DM in patients with TM significantly increases the risk for cardiac complications, heart failure, hyperkinetic arrhythmias and myocardial fibrosis.28

**Growth hormone axis**

Short stature and delay of growth are common features in TM with growth hormone (GH) deficiency being one of the main causes. Treatment of children with TM and reduced GH reserve with recombinant GH has been shown to be effective in increasing significantly linear growth velocity. However, information on treatment of growth hormone deficiency (GHD) in adult thalassaemic patients is lacking.

A study, reassessing the GH - IGF-1 axis, after attainment of final height, in 16 young adults with TM and childhood-onset GHD, found that 19% had persisting GH deficiency.29 For this reason, the GH status should be reassessed in adult TM patients with childhood-onset GHD.

Cross-sectional study on the prevalence of GHD in 94 adult thalassaemic patients with mean age 31.5 years demonstrated severe GHD in 22.3% and partial GHD in 19.1% of patients.30 All patients underwent GHRH plus arginine test; severe GHD was defined by GH peak < 9 mcg/l, while partial GHD was defined as peak GH <9 – 16.5 mcg/l. A similar study of 28 adult thalassaemic patients found that 32.1% had severe GHD, defined as peak GH < 11.0 mcg/l for BMI < 25 and peak GH < 8.0 mcg/l for BMI> 25.31 The lack of correlation between ferritin levels and GH peaks indicates that iron overload is probably not the only determinant in the pathogenesis of GHD in TM. In addition to iron deposition in anterior pituitary, other mechanisms such as cell damage deriving from tissue hypoxia may be relevant. Another similar study found that 25% of adult thalassaemic patients had severe GHD, while 69% of patients had serum IGF-1 values below the 95% confidence interval for age.32

All these studies showed also a very high proportion of patients with IGF-1 deficiency which could not be explained by GHD alone. Noted that GH acts either directly or indirectly through IGF-1. Plausible mechanism are GH resistance attributed to GH receptor alterations or post-receptor defects and reduced liver synthetic activity which contribute to greatly impaired IGF-1 production in patients with TM. In total, a variety of abnormalities has been described in a rage proportion of thalassaemic patients, including classic GHD with low spontaneous GH secretion and impaired response to GH-releasing hormone; GH resistance with reduced IGF-1 response on IGF-1 generation test after GH administration; a combination of classic GHD and GH resistance.33

GH can have an important impact on the quality of life and sense of wellbeing. GH plays an important role in lipid and glucose metabolism and also has anabolic effect on body composition and bone mass. GH
Adrenals

Adrenal insufficiency, manifested as adrenal crisis, appears to be extremely rare in thalassaemic patients. However, several recent studies have reported significant prevalence of “biochemical” adrenal insufficiency, ranging from 18 – 45 %. A recent study, including 56 thalassaemic children and adolescents, showed that 37.5% of participants had adrenal insufficiency confirmed in ITT, the gold standard test.34 This study was unique for two important reasons; first, this was the only study using ITT, considered as the gold standard for the diagnosis of adrenal insufficiency, and second, demonstrated that cortisol binding globulin (CBG) levels were normal, indicating serum total cortisol levels were a good surrogate marker of serum free “active” cortisol. In total, 19.7% of patients had after ITT peak serum cortisol < 14 mcg/dl and they should be strongly considered for regular hydrocortisone replacement, while 17.8% had peak serum cortisol 14-20 mcg/dl and would need steroid cover for major illnesses or operations.34 This study concluded that all thalassaemic patients should have 1 mcg cosyntropin (synthetic ACTH) test as an adrenal function screening test. In the light of 1 mcg cosyntropin test being false positive in 18% of cases, for definite diagnosis, ITT should be performed in those having serum peak cortisol value < 16 mcg/dl after 1 mcg cosyntropin test.34 Another study of 124 adult patients with β-TM major demonstrated a subnormal cortisol response after 1mcg cosyntropin test in 32.1% of patients. It is worth mentioning that only 1.7% of patients would qualify for regular hydrocortisone replacement, while the remaining 30.4% of patients would potentially benefit from steroid cover during major illnesses and preoperatively. Most patients in this cohort exhibited a subtle impairment of adrenocortical function which may become clinically relevant preoperatively. Most patients in this cohort exhibited a subtle impairment of adrenocortical function which may become clinically relevant preoperatively. The majority of patients with clinical hypoparathyroidism have permanent and irreversible.40 Thyroid function should be assessed annually with measurement of serum free T4 and TSH from the age of 9 or earlier if there is a clinical suspicion on hypothyroidism.

Parathyroids

The prevalence of over hypoparathyroidism varies between 1.2 and 19.0%, while subtle subclinical abnormalities in parathyroid hormone (PTH) secretion are much more common with limited clinical significance. The majority of patients with clinical hypoparathyroidism have a mild form of the disease with paraesthesia being the only clinical manifestation. In rare occasions, patients can develop more severe form of hypoparathyroidism and present with tetany and seizures. The variety of clinical manifestations of hypocalcaemia are mainly due to increase of neuromuscular excitability. These include tingling of fingers, toes or lips; numbness of fingers, toes or lips; cramps; carpopedal spasm; stridor due to laryngospasm and seizures. All patients should have annual screening for hypoparathyroidism from the age of 16, including plasma calcium, phosphate and PTH levels. Low plasma calcium levels in combination with high phosphate and low or even inappropriate normal PTH levels indicates hypoparathyroidism.19

Treatment of hypoparathyroidism aims not to achieve normalisation of plasma calcium levels, but to render the patient asymptomatic with a plasma calcium at the lower end of normal range or even just below the normal lower cut-off. The reason for this is that the renal retention of calcium mediated by PTH has been lost. Therefore, any attempt to raise plasma calcium well into the normal range is likely to result in hypercalciuria with the associated risks of renal stones and nephrocalcinosis. Treatment of hypoparathyroidism contains:
- Hydroxylated vitamin D derivatives. Vitamin D requires 1a-hydroxyla-
tion by the kidney to become active. In hypoparathyroidism, there is deficiency of PTH which is essential for 1α-hydroxylation of vitamin D. The hydroxylated vitamin D analogues are either 1α-hydroxycholecalciferol (1,25-dihydroxycholecalciferol) at an initial dose of 0.25 – 0.5 mcg orally once per day or 1,25-dihydroxycholecalciferol (calcitriol) at an initial dose of 0.25 mcg orally twice per day. In the initial stages after initiation of treatment, plasma calcium and phosphate levels should be monitored closely (at least weekly) in order to avoid hypercalcemia. The same applies to every change in the dose of vitamin D analogues with measurement of plasma calcium at least 1 week after. After establishing the appropriate maintenance treatment dose which achieves target calcium at the lower end of normal range, plasma calcium levels should be measured every 3 months. The usual maintenance dose for both vitamin D analogues is 0.25 – 1 mcg/day.

- Oral calcium supplements may be needed in some patients, especially if dietary calcium intake is inadequate. Usual does are 1-3 gr of elemental calcium per day by mouth.

References


