Pregnancy in women with thalassemia: challenges and solutions

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Abstract: Advances in treatment of thalassemia have led to the aging of thalassemic patients, and consequently concern about successful reproductive outcome is augmented. Although women with thalassemia intermedia only were considered competent of achieving pregnancy, case series reveal the willingness of both thalassemia major and thalassemia intermedia women to have a family. Pregnancy in general is characterized by dynamic multiple-system changes and increased susceptibility to oxidative stress, while homozygous, transfusion-dependent, β-thalassemia patients manifest cardiac, hepatic, endocrine, and metabolic disorders attributable to chronic anoxia and iron overload and thalassemia intermedia, usually nontransfused, is associated with augmented risk of thromboembolic events. Pregnancy in thalassemia should be considered a high risk for both mother and fetus, and favorable outcomes are the result of continuous preconception, antenatal, and postpartum assessment and management by a team of thalassemia experts.

Keywords: thalassemia, pregnancy, chelation, transfusion, iron

Introduction

Hemoglobinopathies are among the most common inherited diseases: approximately 7% of the global population is a carrier, and 300,000–500,000 children are born with a severe hemoglobin disorder annually. They are classified according to the impaired globin chains and whether this disorder leads to reduced production of a normal chain or an abnormal tertiary structure of globin chains. The precise structure of the globin chains is coded by genes of chromosomes 16 (the α-gene cluster, comprising the α- and ζ-globin chains) and 11 (the β-gene cluster, comprising the globin chains γ, ε, β, and δ). Hemoglobin must have the correct structure and be trimmed in such a way that the number of α-chains precisely matches that of the β-chains. In adulthood, hemoglobin consists of approximately 98% HbA (α2β2), less than 3% HbA1 (α2δ2), and traces of HbF (α2γ2). The term “thalassemia” refers to hemoglobinopathies characterized by partly or completely suppressed synthesis of one of the two types of polypeptide chains (α or β) as a result of missense/nonsense mutations (single-base substitutions) or frameshift mutations of the genes controlling the structure of the hemoglobin-protein chains in one or both “allelic” globin genes, providing decreased hemoglobin concentration, microcytosis, and anemia. Depending on the genes affected, the resulting defect, and the corresponding effect on the globin chain, several types of thalassemia have been described, the most common types of clinical importance being α-, β/δ-, and β-thalassemia.

Genetic basis and pathophysiology of thalassemias

α-Thalassemia is the most common inherited disorder of hemoglobin, is characterized by reduced or suppressed production of α-globin chains, and occurs at particularly high
frequency in populations from sub-Saharan Africa through the Mediterranean region and Middle East, to the Indian subcontinent and East and Southeast Asia. Almost 5% of the world’s population are carriers, and approximately 1,000,000 patients are affected by various α-thalassemia syndromes worldwide. The α-globin chain synthesis begins in fetal life. The responsible genes – four in total – are situated in two genetic loci on chromosome 16. Gene deletion or less commonly mutation results in α-thalassemia, and phenotype depends on the affected gene number. When all four genes are affected (−/−/−/−) in homozygous α-thalassemia, fetal synthesis of α-chains is impossible, leading to an excess of γ-chains and forming the unstable Bart’s hemoglobin (γ₄), which is incapable of oxygen exchange. The affected fetuses sustain severe anemia, cardiomegaly, and hydrops fetalis, and ultimately intrauterine or neonatal death. When three genes are affected (α⁺/−/−/−), α-chain synthesis is restricted to a minimum. Therefore, β-chains that exist in excess form the unstable HbH (β₄). HbH disease has a phenotypic variability based on mutation type, ranging from mild anemia (deletions on chromosome 16) to a transfusion-dependent one. The existence of two α-genes (α-thalassemia trait) is expressed as a mild hypochromic microcytic anemia. Globin synthesis is still unbalanced, leading to hemolysis and iron overload. In α₀-thalassemia, the two deleted genes belong to the same allele (−/−/α₁/α₁), and this is prevalent among Asian and Eastern Mediterranean populations, while in α⁺-thalassemia, prevalent among African people, the deleted genes belong to different homologous chromosomes. In “silent” carriers, only one α-gene is affected (α⁻/−/α₁/α₁), and the three functional remaining ones are capable of normal hemoglobin production. 

β-Thalassemia is extremely heterogeneous in terms both of genotype and phenotype, depending on the nature of β-gene mutation and the extent of impairment in β-globin chain production. As a rule, heterozygous carriers of β-thalassemia (one affected allele), are asymptomatic, and only altered laboratory values (low, normal, or slightly subnormal hemoglobin levels, slightly low mean cellular hemoglobin, low mean cell volume, low β₃-α₂ globin chain ratio on biosynthesis, HbA₂ ≥3.5%) are observed. In contrast, inheritance of two defective β-globin genes results in a wide phenotype spectrum, ranging from transfusion-dependent (thalassemia major [TM]) to mild or moderate anemia (thalassemia intermedia [TI]). β⁰ refers to the complete absence of production of β-globin on the affected allele, β⁺ refers to alleles with some residual production of β-globin, and β⁺⁺ to a very mild reduction in β-globin production. More than 200 thalassemic mutations have been reported. TI mutations in both parental genes lead to a moderate reduction in β-globin production. TI represents up to a quarter of β-thalassemia patients with a wide spectrum of genotypes and a clinical phenotype ranging between transfusion-dependent thalassemia and the asymptomatic carrier state. Patients have in general later clinical onset, milder anemia not requiring transfusions for survival during the first few years of life, and quality of life is not severely impaired, but the clinical course of the disease, if remaining untreated, is complicated by the multiple effects of chronic hemolytic anemia and the consequent tissue hypoxia, as well as by their compensatory reactions, including increased erythropoiesis with bone marrow expansion and increased intestinal iron absorption.

β-TM or Cooley’s anemia (β⁰/β⁰ or β⁺⁺/β⁺⁺) is characterized by severe hypochromic microcytic anemia, which becomes symptomatic at infancy or early childhood and is apparently transfusion-dependent. The globin chain-synthesis reduction leads to an unbalanced β₃-α₂ globin chain production, where the chains in abundance precipitate, forming erythrocyte inclusions. Pathophysiology is characterized by damaged red blood cells, hemolysis, and erythroid-precursor release in the peripheral circulation, due to ineffective erythropoiesis. The phenotype includes anemia, bone marrow expansion, skeletal deformities, growth restriction, and late sexual maturity. 

As far as it concerns the β-thalassemia population, advances in care by optimal blood transfusion and iron-chelation therapy have improved patient survival into adulthood, as well as quality of life. Consequently, concern about favorable reproductive outcome has increased, as a result of patients’ desire to create their own family. Although women with TI were considered competent of achieving pregnancy after the first observations in the mid-1960s, an increasing number of spontaneous or assisted reproductive technology conceptions in both TI and TM have been reported. 

This article attempts to review available evidence documenting pregnancy in thalassemia, focusing mainly on β-thalassemia and its appropriate follow-up, as it can be a high-risk state for both mother and fetus, and close counseling and monitoring starting from the preconception stage is mandatory.

**Fertility in β-thalassemia**

Hypogonadotropic hypogonadism (HH) is the most frequent endocrinopathy in transfused patients with TM: 51%–66% of thalassemic patients with marked hemosiderosis are predisposed to develop pubertal failure, sexual dysfunction,
infertility, and short stature.25,28 Iron accumulation in the anterior pituitary gland, a tissue with high levels of transferrin receptor, results in free radical oxidative stress, impairing gonadotropins and growth-hormone secretion.27 Furthermore, thalassemias may also suffer from iron accumulation in the ovaries or testes, and oxidative stress may be developed there, when imbalance between the generation of reactive oxygen species and the scavenging capacity of antioxidants in the reproductive tract is present.28 According to the literature, reactive oxygen species may have an important regulatory role through various signal-transduction pathways in the normal functioning of the reproductive system and in female infertility, affecting multiple physiological processes from oocyte maturation to fertilization, embryo development, and pregnancy,29 while recent studies have shown significant acute changes in the hormonal environment and sperm parameters of iron-overloaded patients.30–32 Susceptibility to HH development seems to be associated with genotype, as patients with severe underlying molecular defects have a greater rate of iron loading and probably a different vulnerability to free radical damage.33,34 Furthermore, HH is related to iron toxicity in adipose tissue, impairing and changing the physiological role of leptin – acting as a permissive signal allowing puberty – in sexual maturation and fertility.35,36 The possible role of liver dysfunction and the presence of other endocrinopathies, such as diabetes or hypothyroidism, should not be underestimated when assessing fertility,37 while chronic hypoxia seems to have a crucial role, as studies have revealed decreased gonadotropin secretion within even 2 days of arrival at moderate altitude, enhancing the role of chronic anemia.38

Singer et al suggested that ovarian reserve is preserved in the majority of TM patients less than 30–35 years old, despite a low follicle count and reduced ovarian volume, and that anti-Müllerian hormone could be used as a sensitive marker for ovarian reserve independently of gonadotropin effect correlated with non-transferrin-bound iron, suggesting a role of labile iron in the pathogenesis of decreased reproductive capacity.39 Pulsatile gonadotropin-releasing hormone infusion for ovulation induction is only possible at the early stage of hypothyamic–pituitary damage, but as the majority of patients with HH are nonpulsatile with functional gonads, they are more likely to benefit from human chorionic gonadotropin/human menopausal gonadotropin therapy, which has an 80% success rate.40 Ovulation induction should only be undertaken by a specialist reproductive team, according to Human Fertilisation and Embryology Authority (HFEA) guidelines,41 keeping in mind and making women aware of the risk of hyperstimulation syndrome, multiple pregnancy, ectopic pregnancy, and miscarriage. Around 1%–2% of inducted ovulation cases develop severe hyperstimulation syndrome causing fluid retention with bloating, breathlessness, and nausea, resulting in abdominal pain, vomiting, dyspnea, and rapid weight gain, while the most severe cases have to be hospitalized because of hypovolemic shock, renal and/or respiratory insufficiency, and arterial thromboembolism.42 Protocols forthalassemic women usually involve standard regimes. Patients with endometrial or fallopian tube damage respond better to in vitro fertilization (IVF) programs.

Spermatogenesis in thalassemic males is more difficult, with a success rate of 10%–15% in moderate to severely iron-loaded patients.40 The induction process must be undertaken according to HFEA guidelines, with an emphasis on consent and counseling.41 Micromanipulation techniques, such as intracytoplasmic sperm injection (ICSI), have improved conception rates, even in oligoasthenospermic patients. Sperm should be cryopreserved in all subjects unless azoospermic to preserve fertility and the chance of conception better. However, thalassemic patients with low sperm concentrations are more likely to have a higher degree of defective chromatin packaging, while the negative association between ferritin levels and abnormal sperm morphology suggests a possible detrimental effect on spermatogenesis by the iron chelators.41 Thalassemic patients considered for assisted conception procedures should be counseled accordingly, as mutagenic risk in these individuals, especially after ICSI, where the natural protective barrier against gamete selection during fertilization is lost, seems to be high.44

Prepregnancy planning
Pregnancy planning is essential in both spontaneous and assisted reproductive technology conceptions. Cardiac and liver function, infection control, screening for endocrinological abnormalities, and medication review should be considered regarding females, while both partners should be checked for hemoglobinopathies.45

Screening for thalassemias – genetic counseling
Identifying high-risk populations for thalassemia is the main step for reducing incidence. Screening programs may differ throughout the world, depending on population needs, culture, and/or ethics, and although antenatal diagnosis remains a personal choice, policies are focused on education and counseling.46,47 In Greece, where carriers account for 7.5% of the general population, such a program has been in place since the 1970s, raising awareness and drawing attention to this
inherited disease.48,49 Traditionally, thalassemia was known of higher prevalence in populations of the Middle East, Eastern Mediterranean, India, and Africa, but freedom of movement and subsequent immigration of populations, as well as interethnic mixing, has altered trends. However, according to Hussein et al, there is a lack of randomized trials of preconception genetic risk assessment, and evidence for current policy recommendations is limited to nonrandomized studies.50

Hemoglobin electrophoresis remains the gold standard for the diagnosis and classification of thalassemia. Quantitative evaluation of HbA2 can be made by either electrophoresis or by high-pressure liquid chromatography. Nevertheless, the latter has the additional advantage of quantifying HbF at the same time. Carriers of the β-thalassemia trait demonstrate increased values of HbA2 and HbF.51–53

Where both parents are carriers of the same trait (α–α or β–β couple), genetic counseling should be performed so as to achieve prenatal diagnosis. The couple should be informed of the possibility (25%) of a TM fetus. The diagnosis is made either by chorionic villus sampling or by amniocentesis. Chorionic villus sampling has some advantages, as it can set the diagnosis earlier during the first trimester (11th week), more DNA is obtained by placental biopsy, and it is perhaps safer to perform the placenta than the amniotic cavity. On the contrary, amniocentesis has the drawback of being feasible only after the 16th week. The risk of miscarriage does not differ between these invasive procedures, and is estimated to be less than 1%.54–59

When both parents suffer from a certain hemoglobinopathy, use of donor gametes screened for hemoglobinopathies – preferably donor sperm, as sperm can be more easily available from sperm banks – seems to be the ideal option, while adoption always remain an alternative. If the partner of a homozygous parent is heterozygous, preimplantation genetic diagnosis should be proposed. This involves IVF/ICSI, embryo biopsy, and transfer of healthy embryos. Preimplantation genetic diagnosis can be performed at either the eight-cell stage (cleavage stage) on day 3 after fertilization or at the blastocyst stage on day 5 by biopsy of the trophectoderm cells. Cleavage-stage biopsy is often preferred, as it results in more viable blastocysts and offers more time for genetic testing before embryo transfer. Polymerase chain-reaction techniques (nested, multiplex, fluorescent) have overcome limitations of cleavage-stage biopsy with regard to the small number of cells available for assessment.56–67

Cardiac assessment
Pregnancy-induced cardiovascular changes are the result of the increased metabolic demands of both mother and fetus, and are clinically presented by signs and symptoms resembling heart failure.68 During pregnancy, the heart undergoes important structural changes, such as transient left ventricular hypertrophy, in order to support the demanded increase in functional load, which is as great as 25%–30%. In general, pregnant women tolerate valvular incompetence and uncomplicated left-to-right shunts better than stenosis. Upward displacement of the diaphragm by the enlarging uterus causes the heart to shift to the left and anteriorly, so that the apex beat is moved outward and upward. Furthermore, peripartum cardiomyopathy, a rare condition occurring in the last month of pregnancy or during the 5 postpartum months, is a heart disorder with an unpredictable outcome varying from complete recovery, myocardial sequelae, or aggravation, leading to cardiac transplantation or death. Cardiac function should be assessed by a cardiologist and investigated with electrocardiography (ECG), cardiac echo, and 24-hour Holter monitoring of rhythm in terms of symptoms.68

Despite the increased functional cardiac load during gestation and labor, healthy women do not have cardiac reserve impairment. In contrary, the increased functional load on the heart may cause ventricular failure and pulmonary edema to pregnant women with established or misdiagnosed heart disease and low cardiac reserve. Heart disorders are responsible for about 10% of maternal obstetric deaths, and congenital heart disease may be underestimated. Premature labor can be induced by heart failure, while the risk of maternal or fetal death is strongly associated with maternal New York Heart Association functional classification. Women with certain high-risk disorders (eg, pulmonary hypertension, severe valvular disorders, prior postpartum cardiomyopathy) should be discouraged from pregnancy, as there is high correlation with considerable morbidity and mortality rates.23

Cardiac complications remain the leading cause of death for the thalassemic population. Apart from cardiac iron overload due to regular blood transfusions and delayed or inadequate chelation therapy, thalassemic patients have a greater intestinal iron-absorption capacity than normal individuals, resulting from paradoxical hepcidin suppression from dyserythropoiesis.69 Iron stored in cells, including myocytes, in the form of ferritin, hemosiderin, and free iron, is referred as labile cellular iron. The latter form stimulates the formation of free radicals, provoking cellular injury due to peroxidation damage of membrane lipids and proteins.70 In thalassemic women, iron overload increases the oxidative stress of pregnancy, which peaks by the second trimester of gestation, and can induce great damage to the fetus.71 During pregnancy, basal oxygen consumption is increased and changes in energy substrate use by different organs are
revealed, including the fetoplacental unit. As the initially hypoxic placenta matures and its vascularization develops, it becomes rich in mitochondria, producing locally free radicals and increasing free iron. The impaired function of the mitochondrial respiratory chain affects the heart muscle, leading to decreased cardiac muscular contractility and congestive heart-failure development. In TM, the impaired heart is obliged to maintain a high output through a rigid vascular bed induced by chronic hemolysis, subjected to a continuous state of both volume and pressure overload, rendering the left ventricle more susceptible to decompensation, while the gradually increasing pulmonary vascular resistance seems to lead to the development of pulmonary hypertension, which readily precipitates right-ventricular failure.

Thalassemic pregnant women with normal resting cardiac performance and intensive pregestational chelation therapy usually carry out gestation and delivery successfully. However, it is rather uncertain whether a woman with marginally impaired cardiac performance or with myocardial hemosiderosis, present even in well-chelated patients, will work out the pregnancy stress and the referred hemodynamic changes. Cardiac magnetic resonance imaging (MRI) has been proven of high value with regard to preconception cardiac management. It can accurately define iron overload – underestimated when using only ferritin values or echocardiography – and guide and if needed intensify chelation therapy. The ultimate target is $T_2^*$ measurement.

Liver function

During pregnancy, impairment of liver function can be expected, and may present with subtle findings. Aminotransferases might be slightly elevated, warning of life-threatening processes, such as acute fatty liver of pregnancy or HELLP (hemolysis, elevated liver enzyme levels, low platelet count) syndrome. Thalassemic women are prone to cholelithiasis due to hemolytic anemia and subsequent cholecystitis, while impaired liver function due to iron overload may be present. Preconception liver and biliary tract ultrasound assessment should be performed at a minimum to detect liver cirrhosis, fibrosis, and cholelithiasis, and cholecystectomy should be considered before impregnation. Reduced survival in thalassemia is associated with liver iron concentration above 15 mg/g dry weight. MRI of hepatic iron content using $R_2$ and $R_2^*$ techniques provide accurate and reproducible quantitation of liver iron concentration in thalassemia, and in contrast to liver biopsy, which is invasive and associated with risks, including bleeding and procedure-related pain, is generally more acceptable to patients. A target liver iron concentration of less than 7 mg/g dry weight is recommended prior to conception, and in cases of target excess, intensive preconception chelation should be proposed.

Endocrine complications

Apart from HH, other glands may be affected by tissue hemosiderosis. Diabetes mellitus is frequently diagnosed, resulting from insulin resistance, pancreatic iron overload, genetic factors, and autoimmunity. Those women should be referred to an endocrinologist and have serum fructosamine concentrations kept below 300 nmol/L for at least 3 months before conception. Coexistence of thalassemia and diabetes is an indication of monthly assessment of serum fructosamine concentration. Thyroid function should be assessed and treated before pregnancy, in order to avoid
gestational complications (maternal and perinatal morbidity and mortality), and determined periodically throughout pregnancy.92

Osteoporosis and bone deformity are also present in thalassemic patients, as a result of parathyroid dysfunction, hypogonadism, chelation of calcium by iron-chelation drugs, and vitamin D deficiency. Bone mineral density should be assessed by dual-energy X-ray absorptiometry of the spine or hip for all women with osteopenia or osteoporosis, and supplements should be offered in order to optimize serum vitamin D concentrations.93,94

Medication review

Before pregnancy is actively contemplated or ovulation induced, potentially teratogenic medication needs to be reviewed, including oral hypoglycemic agents, bisphosphonates, and ACE inhibitors. Women on oral chelators (deferasirox [DFX] or deferasirox) are recommended to switch to desferrioxamine [DFO] prior to induction of ovulation/ spermatogenesis.45 Medications that should be discontinued at least 6 months prior to fertility treatment include interferon, ribavirin, and hydroxyurea. Hypothyroid patients receiving thyroid-replacement therapy should receive increased doses to ensure they are euthyroid.

Antepartum management

Pregnant women must be reviewed monthly until the 28th gestational week and every 2 weeks thereafter. Preterm birth related to poor maternal condition, fetal distress, multiple-gestation pregnancies, and placental ischemic disease can complicate gestation. Other commonly met obstetric complications include gestational hypertension, gestational diabetes, placental abruption, urinary tract infection, and renal and gallbladder stones.22,24,55 Women should be screened for gestational diabetes at 16 weeks, and if normal this should be repeated again at 28 weeks. Furthermore, heterozygous thalassemic pregnant women in ethnic groups with high diabetes mellitus incidence should be screened for gestational diabetes.90,91,95,96 Splenomegaly can interfere with the enlargement of the uterus and can be complicated by hypersplenism, necessitating splenectomy during gestation or after delivery.15

Maternal transfusion

Anemia in women with thalassemia deteriorates during pregnancy; however, a proportion of them can remain undiagnosed without screening, as anemia can be of a mild degree or absent. Therefore, their pregnancy will usually be uneventful and normally completed.97,98 Thalassemia per se in combination with gestational anemia (secondary to increased fluid compartment of the body) account partly for different complications of the thalassemic pregnancy, such as fetal intrauterine growth restriction (IUGR) and preterm labor.99,100 Most centers transfuse pregnant women aiming to maintain hemoglobin at the preconception goal (>10 g/dL) to ensure appropriate fetal growth.15,18,22 Despite following this approach, IUGR may be present, suggesting the role of other fetoplacental and maternal factors, while transfusion-acquired red-cell antibodies should be checked prior to pregnancy.101

Referring to TI never- or minimally transfused women, when blood transfusion is necessary, extended genotype and antibody screening should be performed and fully phenotyped matched blood given, as the major fear of inaugural blood transfusions during pregnancy is the developing of alloantibodies, which can stimulate anemia aggravation and grow into severe hemolytic anemia obstructive to transfusions, along with viral transmissions, raising the complication rate. The decision to initiate a transfusion regimen is based on the presence of a worsening maternal anemia or evidence of IUGR, and in this case regular transfusions should be considered. The largest case series of TI pregnancies reported IUGR in 22% of pregnancies.102 However, a recent Italian TI case series with trials of random transfusion regimens administered based on total hemoglobin level but also on general and cardiac maternal status and fetal growth showed that most babies were appropriate for gestational age.22 In general, nontransfused women with hemoglobin ≥8 g/dL at the 36th week of gestation should be advised not to initiate blood transfusions,35 and erythropoietin administration could be an alternative.103,104

Hemolytic alloantibody and erythrocyte-autoantibody development complicates transfusion therapy in thalassemia patients as the rate of red blood-cell alloimmunization following one single blood-unit transfusion is 1%–1.6%, while the rate in patients receiving regular blood transfusions may be as high as 60%.105–107 Alloantibodies crossing placenta can cause fetal and/or neonatal hemolytic anemia, making extended genotype and antibody screening necessary before transfusion, and if transfusion is necessary, fully phenotyped matched blood should be given. Referral to a fetal medicine specialist for consideration of invasive treatment should take place if the middle cerebral artery peak systolic velocity – a noninvasive test for fetal anemia – rises above 1.5-fold the median threshold or if there are other signs of fetal anemia.108

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Chelation during pregnancy

Throughout pregnancy, acceptable levels of maternal hemoglobin should be maintained to minimize hypoxia, but intensive transfusion treatment can aggravate hemosiderosis in patients with a preexisting iron-overload status, elevating oxidative stress and inducing organ failure, whereas women identified with cardiac hemosiderosis or borderline left-ventricle function may bear heart complications.\textsuperscript{109} Therefore, the role of chelation during pregnancy should be clarified.

Chelation therapy will reduce iron overload and help free radical scavenging, reducing the inflammatory process,\textsuperscript{110} but DFO fetotoxicity has not yet been definitely assessed. Although animal studies may identify drug teratogenic effects and skeletal anomalies, the large molecular size and charge of DFO makes placenta crossing doubtful.\textsuperscript{111,112} Various case reports describe its use in early pregnancy,\textsuperscript{21,45,78,113} as well as a large case series of 32 TM women chelated with DFO during the second and third trimesters with favorable fetal outcomes.\textsuperscript{20} According to experts, DFO should be avoided during the first trimester, but subcutaneous administration may be considered in the second and third trimesters for patients with a strong indication for treatment.\textsuperscript{114} As to oral chelators, according to experimental models placental transfer of DFX is minimal in rats,\textsuperscript{115} but fetal outcome data in humans are limited to spontaneous pregnancy case reports,\textsuperscript{116,117} and based on the product label DFX use is contraindicated in pregnant women.

During pregnancy, chelation should be restricted for cases where the potential benefit outweighs the potential fetal risk. If a woman describes cardiac symptoms during pregnancy, she should immediately be assessed by a cardiologist with expertise in thalassemia, in order to balance the risks and decide on the need for chelation treatment.\textsuperscript{78,118,119}

Pregnancy and thromboembolic events

Gestation predisposes to hypercoagulation. Venous thromboembolism frequency is as high as 0.76 to 1.72 per 1,000 gestations, while maternal death is caused prominently by pulmonary embolism.\textsuperscript{120} Platelet activation, fibrin generation, and coagulation factors II, VII, VIII, and X are increased, contrary to fibrinolytic activity and free protein S levels, which are decreased. Additional risk factors for venous thromboembolism during pregnancy and the postpartum period are the reduction in venous flow velocity during gestation, inherited thrombophilias, antiphospholipid syndrome, and previous history of thrombosis.\textsuperscript{121}

Thalassemic women have an increased risk for thrombosis, as the disease entity is a chronic hypercoagulable state with high incidence of thromboembolic episodes, especially in TI, with a risk of as high as 29\%, especially in splenectomized and nontransfused patients.\textsuperscript{122–125} Inherent red-cell defects that activate endothelial cells, creating a procoagulant state, along with platelet abnormalities, deficiency of coagulation inhibitors, cardiac and liver dysfunction, and hormonal deficiencies, seem to comprise the pathophysiology.\textsuperscript{125} Therefore, frequently transfused patients have lower rates of thromboembolism compared with those who are not.\textsuperscript{123,125}

Thrombophylaxis might be essential during pregnancy and the postpartum period in cases of nontransfused TI, splenectomy, or a history of recurrent abortions.\textsuperscript{126} According to recent data, low-dose aspirin, frequently administered to splenectomized β-thalassemia patients, seems to be effective in preventing preeclampsia, preterm birth, and IUGR in high-risk pregnancies without posing a major safety risk to mothers or fetuses.\textsuperscript{127} Therefore, splenectomized women or those with a serum platelet count above 600\times10^{9}/L should begin or continue taking aspirin at a dose of 75 mg/day. Splenectomized women with a platelet count above 600\times10^{9}/L should additionally be offered low-molecular-weight heparin.\textsuperscript{22,126–129}

Fetal monitoring

The first ultrasound scan should be performed at the 7th–9th week of gestation, as these women have a high risk of miscarriage and multiple gestations. In addition to first-trimester (11th–14th weeks) and second trimester (18th–21st weeks) scans, serial fetal biometry scans should be performed monthly after the 24th gestational week, focusing on possible IUGR as a result of chronic maternal anemia and other nutritional elements depletion.\textsuperscript{22,89} The percentage of babies with IUGR varies among different studies as spontaneous conceptions, IVF cases, and singleton and multiple pregnancies are encountered.\textsuperscript{22,33} Miscarriage occurs mostly in cases of thalassemia homozygous fetuses or those with severe IUGR.\textsuperscript{74} The reported frequency varies from 9\% to almost 33\%, which can be explained by gestations conceived spontaneously and through IVF.

Medication modification

Women should be advised to modify their lifestyle and diet, avoid smoking and alcohol, and start taking supplements of folic acid, calcium, and vitamin D. Before and throughout pregnancy, as well as during breast-feeding, adequate calcium and vitamin D intake and bisphosphonate interruption is mandatory, as both are negative calcium-balance states. Especially for thalassemic women, often osteoporotic and vitamin D-deficient, vitamin D levels should be optimized.
before pregnancy and thereafter maintained within the normal range. Folate demand in pregnancy is normally increased, and all thalassemic women are advised to receive folic acid supplementation at a dose of 5 mg/day, in order to prevent fetal neural tube defects, as well as a significant increase in predelivery hemoglobin level, and in heterozygous cases to prevent superimposed megaloblastic anaemia.

Intrapartum management

Time and mode of delivery should be individualized for thalassemia per se, as an uncomplicated disease course should not be considered a proper indication for CS. However, most cases of CS can be attributed to the higher frequency of cephalopelvic disproportion, mainly due to short maternal stature and skeletal deformities combined with normal fetal growth. In case of CS, epidural anesthesia is preferable compared to general anesthesia, as severe maxillofacial deformity in TM patients, especially the older ones, may augment difficulties of intubation. Under this point of view, it is important to correct osteoporosis where required preconception, as spinal abnormalities associated with TM are relevant to regional blockade.

If vaginal delivery is decided, active management of the third stage of delivery is recommended, as this intervention is supposed to reduce blood loss. Fetal hypoxia is common during labor, and thus continuous electronic fetal monitoring is recommended. Transfusion-dependent women that are not chelated will have high concentrations of non-transferrin-bound iron in the serum, a toxic iron form, which can cause cardiac dysrhythmias in combination with labor stress. Therefore, intravenous DFO – 2 g over 24 hours – is recommended for the duration of labor.

Postpartum management

During the postpartum stage, there is a high risk of venous thromboembolism for women with thalassemia, and low-molecular-weight heparin prophylaxis should be administered in hospital, followed by a 7-day postdischarge regimen after vaginal delivery or a 6-week regimen after CS. Women should be referred to a cardiologist after labor, as postpartum cardiac complications have been confirmed. Finally, they should be encouraged to breast-feed, as it is safe in all cases except for those who are HIV, hepatitis C RNA-positive, and/or HBV surface antigen-positive, because of the risk of transmission via breast milk. Significantly lower rates of breast-feeding maintenance when compared to the general population could be explained by the need to restart chelation therapy with oral agents, both of which are contraindicated during breast-feeding, while postpartum chelation using DFO seems to be safe, as DFO is not orally absorbed. Calcium and vitamin D supplements should be continued during breast-feeding, but bisphosphonates should be resumed after cessation of breast-feeding.

Conclusion

Advances in chelation treatment along with regular transfusions have introduced a new era for the thalassemic population, increasing the average life span and rendering the perspective of reproductive capacity attainment and creation of a family, a rational goal for patients and a great task for relevant clinicians. Pregnancy in TM and TI should be considered a high-risk pregnancy, although gestation can be completed safely for both mother and fetus as long as pregnant women follow close screening and are referred to thalassemia specialists. However, as different experts suggest, a standard management plan should be applied to this group of pregnant women, starting with a complete preconception assessment, in order to evaluate and reduce possible risks during pregnancy. Evaluation should include thorough checking for hormonal abnormalities, infectious diseases, liver function, and coagulation status. Women carefully managed and well chelated during preconception usually carry out a successful gestation and labor. Hemoglobin concentration must be maintained over 10 g/dL, and chelation must be stopped as soon as pregnancy is diagnosed. The most important factors to be assessed are cardiac function and iron load, using accurate MRI techniques. In cases of left ventricle-dysfunction development during gestation, and bearing in mind the maternal benefits and the potential fetal risks, DFO could be considered, especially after the critical period of organogenesis. Although CS subtracts from the additional stress of labor, it is usually limited to similar cases as in the general population, and time and mode of delivery should be individualized.

Further studies, as well as national and international registries for thalassemic pregnancies, should be organized and analyzed in order to establish guidelines for this important life period of the thalassemic woman.

Disclosure

The authors report no conflicts of interest in this work.

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