Fanconi Anemia and pregnancy: a new reality
Anemia de Fanconi e gravidez: uma nova realidade

Helena Gomes¹, Inês Lourenço¹, Isabelina Ferreira², Leonor Aboim³, Carlos Veríssimo⁴
Hospital Beatriz Ângelo

Abstract

Fanconi Anemia (FA) is a rare hereditary syndrome which commonly causes infertility. We present a case of a spontaneous successful pregnancy in a patient with FA. During pregnancy, she had a progressive worsening of baseline thrombocytopenia without improvement with corticosteroid or platelet transfusions. Due to inability to maintain a safe platelet level, therapy with a thrombopoietin receptor agonist (Eltrombopag), previously used in patients with refractory immune thrombocytopenia, was started with good response. As this report shows, pregnancy with good perinatal outcome in patients with FA is possible with appropriate multidisciplinary follow-up.

Keywords: Fanconi anemia; Pregnancy; Thrombocytopenia; Thrombopoietin receptor agonist.

CASE DESCRIPTION

We present the case of a 28 years old female patient, nulliparous, diagnosed with FA at 14 years old. Genetic testing showed homozygous presence of variant c.2870G>A in the FANCA gene. Since then, she was followed at the Portuguese Institute of Oncology in Lisbon and had regular laboratory monitoring and performed bone marrow assessments annually. Over the years, laboratory monitoring showed non-progressive findings: hemoglobin ranging 11-13g/dl, average globular volume around 100fl (vitamin B12 and folic acid deficits excluded) and mild thrombocytopenia (platelets 110x10³/μL). Sequential bone marrow trephine biopsy, showed hypocellular marrow without myelodysplasia, leukemia or clonal changes. Other medical specialties performed regular evaluations with normal findings.

Pregnancy developed spontaneously. The first trimester was uneventful, and the blood tests showed mild thrombocytopenia (93x10³/μL platelets). As pregnancy progressed platelet numbers worsened (Figure 1). At 18 weeks, with moderate thrombocytopenia (41x10³/μL platelets), the patient was started on prednisolone 20mg/day.

Second trimester ultrasound didn’t reveal any malformations. At this time platelets declined further to 23x10³/μL. From hereafter, patient’s blood tests were evaluated weekly. Prednisolone dose was further adjusted to 30mg/day, but without any improvement. At 24 weeks, severe thrombocytopenia (13x10³/μL platelets) developed, corticosteroid was discontinued, and platelet transfusions were needed. Despite a good response after the first transfusion (increment to 74x10³/μL platelets), she soon became transfusion dependent, increasing their regularity to biweekly two weeks after.

Given the clinical worsening, the need to optimize the platelet count at delivery, and to avoid regular transfusions with alloimmunization risk, a multidisciplinary team decided, at 27 weeks, to begin Eltrombopag, a thrombopoietin receptor agonist (initially with 25mg, increasing a few days later to 50 mg/day, a dose that was kept). Subsequent weekly blood test evaluation showed that there was a good response to this therapy, maintaining platelet values around 20 x10³/μL. No other transfusions were required until delivery.

In regular laboratory monitoring of the second trimester pregnancy, gestational diabetes was diagnosed, and a probable iatrogenic etiology by regular adminis-
tration of hydrocortisone pre-transfusion can’t be excluded. It was subsequently controlled only with self-monitoring of blood glucose and diet plan.

At 29 weeks a fetal growth restriction (FGR) was diagnosed, and two weeks later the estimated fetal weight was in the 3rd percentile with Doppler changes. The patient was hospitalized and fetal pulmonary maturation begun. Two days later, delivery was planned due to new Doppler worsening. At that time, the patient had 34 x10^3/μL platelets and another platelet transfusion was performed that enabled a value of 71 x10^3/μL platelets before surgery. She ended up receiving 5 units of platelet concentrate during the pregnancy, with no record alloimmunization. At 31 weeks, a caesarean section was performed without complications, and a male infant with 1250g was delivered with an Apgar score of 5/8.

On the first day postpartum platelets were 77x10^3/μL. The puerperium was uneventful. The newborn was admitted to Neonatology due to very low birth weight and respiratory distress syndrome. Later he was discharged without complications or re-hospitalizations.

Three weeks after delivery, platelet numbers were back to baseline (120x10^3/μL). Eltrombopag was maintained up to 6 weeks postpartum. Platelets remained 100-150x10^3/μL after its suspension.

**DISCUSSION**

FA is a rare inherited heterogeneous disease caused by mutations in one of at least 17 different genes, FANCA being the most common (60-65% of the cases). FA proteins’ major function is to maintain genomic stability, through the properly repair of DNA interstrand crosslinks. The disease leads to bone marrow failure, increased sensitivity to cytotoxic therapies and malignancy predisposition.

Patients usually present pancytopenia during the first decade of life but the diagnosis of the disease may not be achieved until adulthood because variable manifestations of the disease are common. Bone marrow failure eventually occurs in the majority of patients, though the time to onset can be quite variable, and in some cases only a single cell line will be involved (typically thrombocytopenia). This reflects our patient’s case who was diagnosed at 14 years old and presented only thrombocytopenia. Severe neutropenia and thrombocytopenia can lead to potentially life-threatening infections and bleeding, usually the cause of death of these patients.

Congenital malformations occur in 60 to 75 percent of patients and are the most frequently presenting features of disease. The most common are skin findings (hyper- or hypopigmentation or café-au-lait spots),
short stature and thumb or other radial abnormalities\textsuperscript{1,3,11}. However, just like our patient, many patients with FA don't manifest classical physical findings of the disease. In fact, about 40% of patients have no major physical anomalies\textsuperscript{7,15}.

FA patients may have a range of endocrine disorders resulting from anatomical disruption of the hypothalamic-pituitary axis during development, more frequently: hypothyroidism, diabetes and dyslipidemia. Infertility and abnormal progression of puberty are frequent\textsuperscript{11,12}. Premature ovarian failure occurs in over 75% of patients. Hematopoietic stem cell transplant is the only cure available in this setting. Other treatments available include medications that target specific signaling pathways, such as the mTOR pathway and ribosomal bypass mechanisms.

During pregnancy, thrombocytopenia is the most common hemostatic change and carries a high risk of unfavorable outcomes: miscarriages, preeclampsia/eclampsia, FGR and fetal, neonatal or maternal death\textsuperscript{9,11}. This is due to physiological mechanisms: increased volume of distribution; increased peripheral destruction by the placenta; and hormonal suppression of bone marrow.

In our patient’s case, her baseline thrombocytopenia due to FA, usually caused by a decrease in platelet production, was worsened due to the reported thrombocytopenic potential of the pregnancy itself. A multidisciplinary team decided to start thrombopoietin receptor agonist, Eltrombopag, in face of a severe thrombocytopenia without corticosteroid response (which excluded a possible association with an autoimmune component), and the need for recurrent transfusions with alloimmunization risk. Eltrombopag is a thrombopoietin receptor agonist that leads to increased platelet production and is a successful treatment used in cases of refractory idiopathic thrombocytopenic purpura (ITP) that has been used before during pregnancy\textsuperscript{13}.

The administration of this drug made it possible to maintain a stable platelet level with no need of recurrent platelet transfusion, reducing the risk of maternal complications resulting from severe thrombocytopenia.

However, this is a difficult decision as, like most drugs, there are no adequate and well-controlled studies of its use in pregnancy (pregnancy category C by the Food and Drug Administration). Animal studies showed evidence of embryo lethality and reduced fetal weights at maternal toxic doses\textsuperscript{13,14}.

Eltrombopag has oral administration and in ITP should be initiated with a 50 mg/day dose that can be increased to 75 mg/day to maintain platelets ≥50 x 10\textsuperscript{3}/L\textsuperscript{14,15}. In this patient, although these numbers were not achieved, the 50 mg/day dosage was kept considering the unknown risk of the drug in pregnancy and the early diagnosis of FGR. Since the drug was started during the third trimester, there was no risk of embryo lethality. Its administration allowed to reduce the need for recurrent transfusions and thus reduce the associated maternal and fetal complications.

Although FGR was a possible adverse effect of the Eltrombopag, we considered it more likely to be primarily due to the underlying maternal pathology, as it was detected early in relation to the drug administration.

Pregnancy in FA patients is a rare event due to reduced average life expectancy and decreased fertility. However, as our work shows, pregnancy is possible, even in non-transplanted women, with proper multidisciplinary medical follow-up essential to manage not only maternal and fetal risks of complications but also the necessary supportive therapy.

Although not yet described in the literature for FA, and further studies are needed, our work suggests that the use of a thrombopoietin receptor agonist in these cases of refractory cytopenia may, as in ITP, contribute to improved obstetric outcomes.

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CONTRIBUTION OF EACH AUTHOR  
Helena Gomes – main author  
Inês Lourenço – help with clinical data collection and associated bibliography search  
Leonor Aboim – help with clinical data collection and associated bibliography search  
Isabelina Conduto – hematological review of the clinical case  
Carlos Verissimo – obstetric review, help with bibliography search  

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ENDEREÇO PARA CORRESPONDÊNCIA  
Helena Gomes  
E-mail: leninha_mhg@hotmail.com  

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