Chronic Massive Fetomaternal Hemorrhage in a newborn from immigrants.
Clinical and organizational implications

Abstract

Fetomaternal hemorrhage (FMH) refers to the entry of fetal blood into the maternal bloodstream before or during delivery. FMH of more than 30 mL occurs with the frequency of about 1/300. Fetal outcomes may be compromised by still births, hydrops fetalis, cardiac complications, and increased rates of postpartum infant death. In most cases, the cause is not identified. Clinical manifestations of FMH depend on the volume of blood lost and the rate that it occurred. We report a case of chronic massive FMH in a newborn of an immigrant mother with a favorable outcome. Medical visits and tests during pregnancy, including ultrasound scans, were not performed. The baby was hemodynamically stable after birth, manifesting only pallor. The complete blood count revealed severe hypochromic anemia (hemoglobin 3.8 g/dl, hematocrit 14.4%) and reticulocytosis (reticulocyte 25.2%). There was no ABO blood type incompatibility and the result of direct Coomb’s test was negative. The Kleihauer-Betke test revealed 5% of fetal erythrocytes in the maternal bloodstream equivalent to 180 mL. The fact that FMH can occur without prior risk factors, and the diagnosis is often postnatal, underscores the importance of heightened medical suspicion particularly in infants born to immigrants where there is often the lack of prenatal visits.

Introduction

Fetomaternal hemorrhage (FMH) refers to the entry of fetal blood into the maternal bloodstream before or during delivery. Most are small-volume blood transfers from fetus to mother of less than 1 mL; conversely, massive FMH of greater than 30 mL occurs in only about 3 of 1000 pregnancies and often has known prior fetal manifestations, such as decreased movement, sinusoidal heart rhythms, peripartum anemia, and fetal anomalies. Most cases of FMH are
idiopathic in origin, most often spontaneous and involve uncomplicated near term pregnancies.4 We report a case of chronic massive FMH, which was proved by Kleihauer-Betke test, in the newborn of immigrants. Some authors have reported in their diagnosis of FMH the existence of socioeconomic and racial disparity. The purpose of this study is to describe the clinical aspects of the disease but also to emphasize the problems, increasing frequent in Italy, of the lack of visits of immigrant women during pregnancy resulting in the potential delay or missed diagnosis in the newborn.

Case Report

A 23-year-old, gravida 2, para 1, albanian woman at 37 2/7 weeks estimated gestational age delivered a 2980-g infant via elective caesarean section for premature rupture of membranes. Tests during pregnancy, including ultrasound scans, were not been performed. Apgar scores were 7, 8 and 8 at 1, 5 and 10 minutes, respectively. The baby was breathing spontaneously and alert, but extremely pale. Thirty minutes after the birth, the newborn was still pale with systolic murmur 1-2/6 a mild hypotonia and had the following vital parameters: heart rate 168/min, respiratory rate 54/min, normal arterial oxygen saturation and blood pressure 55/33 mmHg (mean arterial pressure 42 mmHg). The pH was 7.34 with a base deficit of 4.5 meq/l. The haemoglobin was 3.8 g/dl, hematocrit of 14.4%, WBCs 13200/mm3 and platelets 230000/mm3. On the peripheral smear, reticulocytes were 25.2% and erythroblasts 28%. Coombs test was negative. The newborn needed fluid therapy and one packed red blood cells transfusion of 20 ml/kg, which raised the hemoglobin level to 10.7 g/dl with a hematocrit of 34.5%. Hemolytic diseases and other causes of perinatal hemorrhage were excluded. The Kleihauer-Betke test revealed 5% of fetal erythrocytes in the maternal bloodstream equivalent to 180 ml according to the Cunningham formula. The newborn improved with the fluid therapy and blood transfusion and was discharged on the 12th day, when a complete blood count showed: hemoglobin 9.2 g/dl, hematocrit 27.5%, reticulocytes 1.5%, WBCs 10600/mm3, and platelets 194000/mm3. Physical and neurological evaluations were normal when the infant was discharged. The cranial ultrasound at day 1 of life showed a bulky choroid plexus on the left side. There was no evidence of periventricular leukomalacia. Follow-up scans on day 7 confirmed a grade I intraventricular haemorrhage (IVH) on the left side. On day 30, mild ventriculomegaly was noted with resolution of the IVH.

Discussion

Fetal blood likely enters the maternal bloodstream during all pregnancies, without apparent clinical significance in most cases. In 96% of the cases, the loss of blood is minimal, usually only less than 0.5 mL and less than 15 mL in more than 99%. FMH of a significant volume (>30 mL) is rare with a frequency of about 3/1000 births.2,3 Thresholds of 80 ml or 150 ml also have been proposed to define “large” or “massive” FMH with an incidence of 0.9/10,000 births and 0.2/10,000 births at cutoffs of more than 80 ml and more than 150 ml, respectively.4 Fetal outcomes may be compromised by still births, hydrops fetalis, cardiac complications, and increased rates of postpartum infant death.2 The risk factors of FMH include antepartum fetal death, cesarean delivery, abruptio placenta, placenta previa, manual removal of the placenta, intrapartum manipulation, antepartum genital bleeding, third-trimester trauma, placental tumors and third-trimester aminocentesis.2,4 More than 80% of cases in which the FMH was estimated to be greater than 30 ml remain unexplained.4 The clinical manifestations and prognosis of FMH depend on the volume of the hemorrhage and the rapidity with which it occurs. In cases of prolonged or repeated bleeding during the course of the pregnancy, anemia slowly develops, giving the fetus the possibility to develop hemodynamic compensation with increased hemopoietic activity (increased reticulocytes and erythroblasts in the peripheral smear).3 The diagnosis is often postnatal and these infants may manifest only pallor at birth.5 In acute FMH, rapid blood loss is followed by perinatal hypoxia and intrapartum death or severe anemia and hypoxia at birth.6 A decrease in fetal movements associated with abnormal cardiotocographic findings, such as a sinusoidal pattern of the fetal heart rate, traditionally equated with fetal anemia, may be a warning sign of a massive FMH, especially in a low-risk pregnancy.5,6 The Kleihauer-Betke (KB) test is the standard method of detecting FMH.7 Comparison with other more expensive or technologically advanced methods such as flow cytometry has shown that the KB stain, like the more advanced methods, is almost equally sensitive in its detection of FMH.8 The red cell morphologic study of this infant indicated hypochromic anemia with an elevated reticulocyte count of 25.2%. Compared with the value of 3-7% in normal newborns, this was markedly elevated. This finding suggested that the course of blood loss was a chronic process, because two to three days are usually required for a reticulocyte response to be observed, with a peak response after 10 to 14 days.9 The estimated volume of fetal blood in the maternal bloodstream was approximately 180 ml according to the Cunningham formula.10 Although a great amount of blood was lost, the patient only demonstrated pallor which indicated a chronic FMH with good compensation. The short- and long-term prognosis for fetuses that experience FMH is variable. A FMH of 20 ml/kg, which represents 20% of the fetoplacental blood volume, is considered massive because it is associated with significant fetal/neonatal morbidity or mortality.1 At this cutoff, 26.1% of the fetuses were stillborn, 17.4% required preterm delivery, 34.8% were admitted to the neonatal intensive care unit, and 21.7% received postnatal transfusions.2 These risks increased with increasing fetal volume transfused. With bleeds greater than 80 ml/kg, two thirds of the fetuses died before deliv-
Cerebral palsy has been linked with FMH, but the likelihood of this occurring among survivors is not well defined. Some authors have shown that the diagnosis of FMH is associated with significant morbidity as well as regional, socioeconomic and racial disparity. These authors have shown that higher patient income was associated with increased likelihood of FMH diagnosis (OR 1.2), and whites were more likely to be diagnosed than ethnic minorities (OR 1.9). As the immigrant birth rate in Italy continues to rise, this becomes increasingly important. According to The National Institute for Statistics, births attributable to foreign mothers are equal to 18.8% of the total in 2010, with peaks of 29% in some regions, compared to 6.4% in 2000. Of these mothers, only 4.8% have an Italian partner while the majority (14%) have a foreign partner. The majority of pregnant immigrants do not perform routine obstetric visits because of religious reasons, linguistic difficulties, and their precarious socio-sanitary and economic conditions. These social and economic hardships are associated with more frequent adverse outcomes for maternal and neonatal health. In particular, there is a higher prevalence of low birth weight babies and perinatal and infant morbidity.

**Conclusion**

The fact that FMH can occur without prior risk factors and the diagnosis is often postnatal underscores the importance of a high index of suspicion particularly in infants born to immigrants where there is the lack of prenatal visits. The case described, first in Italy, highlights the need for particular attention by national authorities to pregnancy health care issues of immigrants.

**References**