Perinatal Outcome of Pregnancies Complicated by Immune Thrombocytopenia

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Abstract

Background: Immune thrombocytopenia (ITP) is an autoimmune disorder that leads to premature destruction of antibody-coated platelets. This study evaluated perinatal outcome and medications used for pregnancies complicated by ITP.

Methods: Medical records of 132 pregnancies belonged to 125 parturients with ITP who delivered between March 2001 and January 2011 were reviewed. Cases were included if diagnosed before pregnancy or if their platelet counts (PCs) were less than 80,000/µL during pregnancy without any other cause. Maternal and fetal outcomes were compared.

Results: Fifty six mothers (42.1%) had PC<50,000, 18 women (13.5%) developed preeclampsia and 15 (11.3%) were diabetics. Corticosteroid was used for 120 cases (90.9%) and intravenous immunoglobulin for 14 women (10.5%). PCs of 114 neonates were available in the charts and 84 (83.2%) had PC>150,000/µL. Three neonates (2.3%) had PC<50,000, 31 neonates (23.3%) had preterm births and 32 (24.1%) needed NICU admissions. Fifty seven cases of ITP (43.2%) were diagnosed before pregnancy and 75 (56.8%) were diagnosed during pregnancy. There were 2 intrauterine fetal deaths and higher NICU admissions, 20 (34.48%) versus 12 (16%) in the first group (p=0.01).

Conclusion: Perinatal outcome of pregnancies with ITP is generally good. However neonates born from parturients with chronic ITP needed more NICU admissions.

Keywords: Immune thrombocytopenia; Perinatal; Outcome; Iran

Introduction

Immune thrombocytopenia (ITP) is an autoimmune disorder that leads to premature destruction of antibody-coated platelets. For the diagnosis of ITP there is no approved confirmatory laboratory test at present. Diagnosis is made by demonstrating an isolated thrombocytopenia without an obvious cause. Antiplatelet antibodies may cross the placenta and cause bleeding complications in the neonate.1,2 Pregnancy with ITP is sometimes difficult to manage due to bleeding potentials for the mother and her fetus during peripartum periods. Thrombocytopenia occurs in 7-10% of unselected pregnancies. However, ITP accounts for only 3% of these cases compared to gestational thrombocytopenia and hypertensive disorders of pregnancy that account for 74% and 21% of the cases, respectively.3,4 This project was designed to retrospectively study the outcome of pregnant women with ITP and their neonates during an almost 10-year period.

Materials and Methods

This study was performed retrospectively from March 2001 to January 2011 in Hafez and Hazrate Zeinab hospitals affiliated to Shiraz University of Medical Sciences, Shiraz, Southern Iran. Thrombocytopenia...
Pregnancies with thrombocytopenia

was defined as platelet count (PC) of less than 150,000/µL. Patients were included if they were known cases of chronic ITP prior to pregnancy or were diagnosed during pregnancy with PCs of less than 80,000/µL for which other causes were excluded.5,6 Other causes of thrombocytopenia include sepsis, HELLP (hemolysis, elevated liver enzymes and low platelet) syndrome, disseminated intravascular coagulation (DIC), drug induced thrombocytopenia, and thrombocytopenia associated with autoimmune diseases such as systemic lupus erythematosis, thrombotic thrombocytopenia, hemolytic uremic syndrome and hereditary forms of thrombocytopenia.5

Incidental or gestational thrombocytopenia was diagnosed when thrombocytopenia was first diagnosed during pregnancy in an otherwise asymptomatic patient with platelet counts of more than 80,000/µL.5,6 These cases were excluded from this study. According to the PCs, the mothers and their neonates were classified to have mild (100,000 ≤ PC < 150,000/µL) moderate (50,000 ≤ PC < 100,000/µL) and severe (PC < 50,000/µL) thrombocytopenia.

A woman was considered to have preeclampsia if her blood pressure was ≥140/90 with urine protein of more than 300 mg in 24 hours or more than 1+ dipstick.9 Women with overt or gestational diabetes who were diagnosed according to ACOG criteria10 were analyzed as diabetic group. If delivery occurred before 259 days of gestational age, it was considered as a preterm birth (PTB).11 Gestational ages were calculated from the first day of the last menstrual period and were confirmed by first trimester ultrasound scans.

The extracted data included maternal PC at delivery, history of ITP prior to pregnancy, prior splenectomy, therapeutic modalities, mode of delivery and complications, neonatal PC at birth, gestational age at delivery, neonatal birth weight and NICU admission. This project was approved by the Ethics Committee of Shiraz University of Medical Sciences. Statistical analyses were performed by SPSS software (Version 16, Chicago, IL, USA). Evaluation of correlation between maternal and neonatal PCs was performed by Pearson correlation test. Evaluation of correlation between previous splenectomy in the mother and neonatal PC was performed by independent sample t-test. Chi-Square test was used for comparison of ratios that were presented as number (%) and for comparison of means, independent sample t test was used. Statistical significance was set at p<0.05.

Results

One hundreds and thirty two pregnancies were included from 125 women with ITP, 7 women had two deliveries and one was twin. Four singleton women did not deliver in our centers, so 129 neonates remained for the final analysis. Fifty seven pregnancies from 132 (43.2%) were known cases of ITP before pregnancy and 75 (56.8%) were diagnosed during pregnancy. The mean gestational age at delivery was 266.9±14.7 days (range: 206-294 days). Eighteen women (13.5%) were considered to have preeclampsia and 15 (11.3%) were diabetic. Several maternal and neonatal data were presented in Table 1.

There were 9 women (6.8%) with PC>150,000/µL. Eight mothers (6%) had mild, 59 (44.4%) had moderate and 56 (42.1%) had severe thrombocytopenia. The platelet counts of 114 neonates were available in the charts. There were 84 neonates (83.2%) with PC≥150,000/µL. Fourteen neonates (10.5%) had mild, 13 (9.8%) moderate and 3 (2.3%) severe thrombocytopenia, respectively. The 3 neonates with PC<50,000/µL were born from the mothers with ITP prior to pregnancy. No neonates had severe bleeding complications and no intracranial hemorrhage occurred.

There was no significant correlation between maternal platelet count and neonatal platelet count (p=0.19, r=0.15). Also, there was no statistically significant correlation between previous splenectomy and neonatal platelet counts at birth (p=0.64).

Thirty-one (23.3%) neonates had PTB and 32 (24.1%) infants had NICU admissions. Eight neonates (6%) had Apgar Scores below 7 at 5 minutes. There were 2 fetuses that had intrauterine deaths on admission time and were included among the cases with Apgar scores below 7. Seventy four pregnancies

| Table 1: Several maternal and neonatal features of pregnancies complicated by ITP at the time of delivery. |
|----------------------------------------------------------|----------------------------------------------------------|
| **Variable** | **mean±SD** | **Range** |
| Maternal age (years) | 26.5±4.8 | 16-40 |
| Maternal platelet count at delivery (/µL) | 62,260±38,642 | 3,000-200,000 |
| Gestational age at delivery (days) | 266.9±14.7 | 206-294 |
| Neonatal birth weight (grams) | 29189±588.36 | 1150-3900 |
| Neonatal platelet count (/µL) | 146,040±60,995 | 36,000-314,000 |
(55.6%) were terminated by cesarean and 54 (40.6%) had vaginal deliveries. Indications for abdominal deliveries were fetal distress (n=32, 43.2%), previous cesareans (n=27, 36.5%), abnormal presentations (n=6, 8.1%), active phase arrest (n=4, 5.4%), twin (n=1, 1.4%), prolonged rupture of membranes and not responsive to induction of labor (n=4, 5.4%).

Medications that were used for the study group during the admission time were corticosteroids including prednisolone (n=88, 66.2%), dexamethasone (n=78, 58.6%) and hydrocortisone (n=39, 29.3%). Fourteen women (10.5%) used intravenous immunoglobulin (IVIg), 38 women (28.6%) used at least one kind of medication, 66 (49.6%) used two kinds, 15 (11.3%) used three kinds and for one case (0.8%), 4 types of medications were used simultaneously. There were only 11 women (8.3%) who did not use any kind of medications. Four of them were diagnosed before pregnancy and 7 cases diagnosed during pregnancy. There were 8 (6%) women known cases of ITP who had splenectomy prior to their pregnancies. Sixty three women (47.4%) received platelet transfusion in their hospital course.

There were 57 (43.2%) cases who were known cases of ITP before pregnancy and 75 (56.8%) were diagnosed during pregnancy. These two groups were compared. The data was presented in Table 2. The two groups had no statistically significant difference except for NICU admission rate which was higher for the neonates of the women who were known cases of ITP prior to pregnancy (p=0.01).

There were two women admitted with intrauterine dead fetuses at 224 and 247 days of pregnancy. They were both known cases of ITP prior to pregnancy. In this study, there were 5 women with bleeding complications who needed blood transfusions. The first was a 28 years old woman with PC=20,000/µL who needed emergency cesarean for fetal distress and developed hematoma at the site of spinal needle. The second case had normal delivery with PC=41,000/µL and developed postpartum uterine bleeding and needed transfusion. The other 3 cases had cesarean deliveries for obstetric indications with PCs of 15,000/µL, 45,000/µL and 70,000/µL and developed hemorrhage during operation. The parturient with platelet count of 70,000/µL needed reoperation for drainage of massive hematomas. There were no maternal mortalities among this study population.

### Table 2: Comparison of maternal and neonatal characteristics of pregnancies complicated by ITP between the cases that were diagnosed before pregnancy with the cases diagnosed during pregnancy.

| Variable                      | Diagnosed before pregnancy (n=58) | Diagnosed during pregnancy (n=75) | P value<sup>c</sup> |
|-------------------------------|-----------------------------------|-----------------------------------|---------------------|
| Maternal age (year)           | 26.08±5.09                        | 26.8±4.64                        | 0.416               |
| Maternal PCs (/µL)            | 64,035±44,941                     | 60,920±33,327                    | 0.648               |
| Gestational age at delivery (days) | 264±17.3                         | 267±14.3                        | 0.204               |
| Preterm birth                 | 16 (27.6%)                        | 15 (20%)                         | 0.416               |
| Diabetes                      | 9 (15.5%)                         | 6 (8%)                           | 0.165               |
| Preeclampsia                  | 7 (12.1%)                         | 11 (14.7%)                      | 0.695               |
| Cesarean delivery             | 35 (60.3%)                        | 39 (52%)                        | 0.348               |
| 5 minutes apgar score<7       | 3 (5.2%)                          | 5 (6.7%)                        | 0.715               |
| Neonatal birth weight (grams) | 2915.7±678                        | 2926.6±526.3                    | 0.924               |
| Neonatal PC<sup>a</sup> (/µL) | 135880±58899                      | 153860±62605                     | 0.924               |
| Neonatal PC<sup>a</sup><50,000/µL | 3 (2.3%)                            | 0 (0.0%)                           | 0.122               |
| Maternal PC<sup>a</sup><50,000/µL | 25 (43.1%)                           | 31 (41.3%)                        | 0.447               |
| NICU<sup>b</sup> admission    | 20 (34.48%)                       | 12 (16%)                        | 0.017               |

<sup>a</sup>PC: Platelet Count; <sup>b</sup>NICU: Neonatal Intensive Care Unit. <sup>c</sup>Values are presented as mean±SD or number (%).

### Discussion

ITP is estimated to occur in 1 in 1000 to 1 in 10 000 pregnant women and is responsible for almost 3% of all cases of thrombocytopenia in pregnancy and considered as a rare condition.<sup>6,12</sup> Careful management of ITP during pregnancy may become life saving for the mother and her offspring. Diagnosis of ITP is a diagnosis of exclusion and should be performed clinically. According to practice guidelines platelets, rarely decrease to less than 80,000/µL in gestational thrombocytopenia.<sup>5,a</sup> In this study, if thrombocytopenia presented for the first time during pregnancy, we included only the cases with PCs less than 80,000/µL. However the cases that were diagnosed prior to pregnancy were all included. We retrospectively reviewed
the admission charts of the patients in an almost 10-year period and 132 pregnancies belonged to 125 mothers were included. Fifty-eight cases were diagnosed before pregnancy and 75 were diagnosed during pregnancy.

The maternal ages ranged between 16-40 years (mean 26.49±4.8 years) and maternal platelet counts were between 3,000-200,000/µL (mean 62,850±39,081/µL). However, 115 women (86.6%) were classified to have moderate to severe thrombocytopenia with platelet counts of below 100,000/µL and 56 of them (42.1%) had PC< 50,000/µL. On the contrast, among the neonates with the mean platelet count of 146,040±60,995/µL, one hundred and thirteen (84.9%) had platelet counts of more than 100,000/µL. Thirteen neonates (9.8%) had platelet counts between 50,000-100,000/µL and only 3 (2.3%) had platelet counts below 50,000/µL. None of these neonates showed severe bleeding complications and there was no case of intracranial hemorrhage. However, higher rates had been reported for severe neonatal thrombocytopenia in previous studies. Neonatal PC may drop during few days after birth and the difference may be caused by measuring PCs at different durations from delivery. This low rate of fetal severe thrombocytopenia emphasizes that performing invasive procedures before delivery for the measurement of fetal PCs may not be justified. Analysis of our data showed no correlation between maternal and neonatal platelet counts.

Although the mean gestational age at delivery was 266.4±15.6 days in this study, 31 neonates (23.3%) had PTBs and 32 neonates had NICU admissions mostly due to prematurity and its complications. The incidence of PTB in most developed countries is about 5-10%. However, higher PTB rates have been reported for some populations, but it is estimated to be about 5% in the general population in our centers. The calculated PTB rate of 23.3% for the pregnancies complicated by ITP in this study was four times higher.

Among this study group, there were 18 women (13.6%) who developed preeclampsia. According to the fact that hypertensive disorders complicate 5-10% of pregnancies, the calculated rate of 13.6% for the parturients with ITP is more than general population. In this study, there were 15 women (11.4%) diagnosed to have either impaired glucose tolerance test or overt diabetes. Compared to the rate of diabetes estimated to be about 4.2%, this study showed a higher tendency for impaired glucose metabolism in this population. Higher rates of preeclampsia and diabetes in this group may be consequences of associated vascular problems or as a complication of corticosteroid therapy.

Antiplatelet antibodies may cross the placenta and cause thrombocytopenia and bleeding complications in the neonate. Obstetricians are always worried about birth trauma and especially ICH in a thrombocytopenic neonate. From a historical point of view, it was suggested to perform umbilical blood sampling near term and cesarean delivery was considered to be indicated for the thrombocytopenic fetuses or for the mothers who have positive circulating platelet antibody test. However evidence showed that invasive tests have little effect in preventing neonatal complications and it is now a general agreement that vaginal delivery is preferred for the parturients with ITP unless an obstetric indication occurs. In this study, 55 women (41.4%) had normal vaginal deliveries and 74 cases (55.6%) delivered abdominally. The most frequent indication for cesareans was fetal distress that occurred for 32 cases (43.2%).

The mean neonatal birth weight was 29189±588.36 grams. Five minutes apgar scores below 7 was seen only in 8 neonates (6%) including 2 dead fetuses. These two fetuses that were dead on admission in addition to the three neonates that had platelet counts less than 50,000/µL were all born from the mothers with chronic ITP. So, we sub-classified all of the women to two groups (diagnosed before pregnancy and diagnosed during pregnancy) and compared the outcomes between them. We noticed that maternal age, maternal platelet counts, gestational age at delivery, 5 minutes apgar scores<7, rate of abdominal deliveries, mean neonatal birth weights and the mean neonatal platelet counts were not statistically different between the two groups. However, NICU admissions were significantly higher in the group that were diagnosed before pregnancy, 20 (34.48%) compared to 12 (16%) (p=0.01). Suggesting that fetus in a chronic case may probably be more compromised and fetal well being evaluations are more strongly recommended for this group during pregnancy.

Eleven women (8.3%) did not receive any medication. However, the most frequently used medication were corticosteroids (n=121, 91.7%). Fourteen women (10.5%) who did not respond to corticosteroids to rise their platelet counts received IVIG during pregnancy. Sixty three (47.4%) had platelet transfusions at the time of normal delivery or cesarean. Some patients in this study were over-treated compared to the clinical guidelines, which reflects the fear of the
medical team from low PCs and its probable complications during delivery. According to the clinical practice guidelines PC>20,000 during pregnancy and PC>50,000 near term and PC>80,000 for epidural anesthesia is targeted. However, counts may drop unpredictably and major bleeding has been reported rarely even with platelet counts that seem appropriate. Furthermore obstetric hemorrhage may happen due to placental abruption or previa, uterine atonia or uncontrollable genital lacerations even in a normal coagulating condition. In this study, there were two cases that had platelet counts above the targeted values but developed bleeding complications. The first case developed severe vaginal bleeding after normal delivery with PC=41,000/µL and the second case had severe hemorrhage in abdominal incision site with PC=70,000/µL. However, the final maternal and neonatal outcomes of pregnancies complicated by ITP are generally good. All of the cases should be carefully managed by a team of hematologist, neonatologist, anesthesiologist and obstetrician for the best results. However, fetal well being evaluations are more strongly recommended for the chronic cases.

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References

1 Cunningham FG, Lenovo KJ, Bloom SL, Haut JC, Rouse DJ, Spong CY. Preterm birth. In: Williams Obstetrics, 23rd Ed. New York: Mc Graw-Hill; 2010: pp. 1079-1103.
2 Neunert C, Lim W, Crowther M, Cohen A, Solberg L Jr, Crowther MA. American Society of Hematology. The American Society of Hematology 2011 evidence-based practice guideline for immune thrombocytopenia. Blood 2011; 117: 4190-207. [21325604] [doi.org/10.1182/blood-2010-08-302984]
3 Karim R, Sacher RA. Thrombocytopenia in pregnancy. Curr Hematol Rep 2004; 3:128-33. [14965489]
4 Clinical practice guidelines August 2006 MOH/pak/115.06 (GU). Management of immune thrombocytopenic purpura. Ministry of Health Malaysia, Academy of Medicine.
5 George JN, Woolf SH, Raskob GE, Wasser JS, Aleford LM, Ballem PJ, Blanchette VS, Bussel JB, Cines DB, Kelton JG, Lichtin AE, McMillan R, Okerblom JA, Regan DH, Warrier I. Idiopathic thrombocytopenic purpura: an explicit method developed by American Society of Hematology. Blood 1996; 88:3-40. [8704187]
6 British Committee for Standards in Haematology. General Haematology Task Force. Guidelines for the investigation and management of idiopathic thrombocytopenia in adults. Children and in pregnancy. Br J Haematol 2003; 120:574-96. [12588344] [doi.org/10.1046/j.1365-2141.2003.04131.x]
7 Rodeghiero F, Slavi R, Gernsheimer T, Michel M, Provant D, Arnold DM, Bussel JB, Cines DB, Cooper N, Godeau B, Lechner K, Mazzucconi MG, McMillan R, Sanz MA, Imbach P, Blanchette V, Kühne T, Ruggen M, George JN. Standardization of terminology, definitions and outcome criteria in immune thrombocytopenic purpura of adults and children: report from an international working group. Blood 2009; 113:2386-93. [19005 182] [doi.org/10.1182/blood-2008-07-162503]
8 Gernsheimer T, McCare KR. Immune thrombocytopenic purpura in pregnancy. Curr Opin Hematol 2007; 14:574-80. [17934366] [doi.org/10.1097/MOH.0b013e3282f6dc2]
9 ACOG Committee on Practice Bulletins—Obstetrics. ACOG practice bulletin. Diagnosis and management of preeclampsia and eclampsia. Number 33, January 2002. Obstet Gynecol 2002; 99:159-67. [16175681]
10 Cunningham FG, Lenovo KJ, Bloom SL, Haut JC, Rouse DJ, Spong CY. Diabetes. In: Williams Obstetrics. New York: Mc Graw-Hill, 2010: pp. 1104-1125.
11 ACOG Committee on Practice Bulletins—Obstetrics. ACOG practice bulletin. Management of preterm labor. Number 43, May 2003. Int J Gynaecol Obstet 2003; 82:127-35. [12834934] [doi.org/10.1016/S0020-7292(03)00247-9]
12 Gill KK, Kelton JG. Management of idiopathic thrombocytopenic purpura in pregnancy. Semin Hematol 2000; 37:275-89. [10942222]
13 Webert KE, Mittal R, Sigouin C, Heddle NM, Kelton JG. A retrospective 11-year analysis of obstetric patients with idiopathic thrombocytopenic purpura. Blood 2003; 102:4306-11. [12947011]
Svigos JM, Robinson JS, Vigneswarn R. Threatened and actual preterm labor including mode of delivery. In: James DK, Weiner CP, Steer PJ, Gonik B, editors. High risk pregnancy: management options. Philadelphia: Elsevier Saunders, 2006: pp. 1304-1320.

Ananth CV, Liu S, Joseph KS, Kramer MS. Fetal and Infant Health Study Group of the Canadian Perinatal Surveillance System. A comparison of foetal and infant mortality in the United States. Int J Epidemiol 2009;38:480-9. [18806 278] [doi.org/10.1093/ije/dyn194]

Namavar Jahromi B, Salarian L, Shiravani Z. Maternal risk factors and neonatal outcome of the admitted patients for preterm spontaneous uterine contractions. Iran Red Crescent Med J 2011;13:877-83. [22737433]

Baraban E, McCoy L, Simon P. Increasing prevalence of gestational diabetes and pregnancy-related hypertension in Los Angeles country, California, 1991-2003. Prev Chronic Dis 2008;5:A77. [18558027]

Kaplan C, Daffos F, Forestier F, Tertian G, Catherine N, Pons JC, Tchernia G. Fetal platelet counts in thrombocytopenic pregnancy. Lancet 1990;336:979-82. [1977013] [doi.org/10.1016/0140-6736(90)92430-P]

Samuels P, Bussel JB, Braitman LE, Tomaski A, Druzin ML, Mennuti MT. Cines DB. Estimation of the risk of thrombocytopenia in the offspring of pregnant women with presumed immune thrombocytopenic purpura. N Engl J Med 1990;323:229-35. [2366833] [doi.org/10.1056/NEJM199007263230404]

Silver RM, Branch DW, Scott JR. Maternal thrombocytopenia in pregnancy: Time for a reassessment. Am J Obstet Gynecol 1995;173:479-82. [7645624] [doi.org/10.1002/0002-9378(199507)173:4<479::AID-OBGYN10>3.0.CO;2-2]

Levy JA, Murphy LD. Thrombocytopenia in pregnancy. J Am Board Fam Pract 2002;15:462-7. [12150462]

Payne SD, Resnik R, Moore TR, Hedrana HL, Kelly TF. Maternal characteristics and risk of severe neonatal thrombocytopenia and intracranial hemorrhage in pregnancies complicated by autoimmune thrombocytopenia. Am J Obstet Gynecol 1997;177:149-55. [9240599] [doi.org/10.1016/S0002-9378(97)00454-X]

Stasi R, Provan D. Management of immune thrombocytopenic purpura in adults. Mayo Clin Proc 2004;79:504-22. [15065616] [doi.org/10.4065/79.4.504]

Provan D, Stasi R, Newland AC, Blanchette VS, Bolton-Maggs P, Bussel JB, Chong BH, Cines DB, Gernsheimer TB, Godeau B, Grainger J, Greer I, Hunt BJ, Imbach PA, Lyons G, McMillan R, Rodeghiero F, Sanz MA, Tarantino M, Watson S, Young J, Kuter DJ. International consensus report on the intervention and management of primary immune thrombocytopenia. Blood 2010;115:168-86. [19846889] [doi.org/10.1182/blood-2009-06-225565]

Cines DB, Bussel JB. How I treat idiopathic thrombocytopenic purpura (ITP). Blood 2005;106:2244-51. [15941913] [doi.org/10.1182/blood-2004-12-4598]