Case report of aplastic anaemia detected in third trimester of pregnancy: dilemmas faced

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ABSTRACT
Aplastic anaemia with pregnancy is rarely encountered. Management of aplastic anaemia in pregnancy primarily involves a multidisciplinary approach offering supportive care. Our case was challenging as she developed aplastic anaemia during the third trimester and had refractory thrombocytopenia. She required platelet transfusions on a daily basis for few weeks as well as packed red blood cells frequently. Her leucocyte count was low initially but improved quickly unlike the platelet counts. Initiation of immunosuppressive therapy turned out to be beneficial and culminated in a good outcome. After starting immunosuppressive therapy with eltrombopag and cyclosporine she drifted through term and achieved a normal vaginal delivery.

Keywords: Aplastic anaemia, Pregnancy, Eltrombopag

INTRODUCTION
Aplastic anaemia is a rare disorder. The incidence of aplastic anaemia in Asia is 4-6 per million which is higher compared to about 2 million in Western countries.1 It is defined as pancytopenia with a hypocellular bone marrow. Most patients with aplastic anaemia present with anaemia and thrombocytopenia. Causes include exposure to chemicals, drugs and viral infections. Treatment options include bone marrow transplantation, antithymocyte globulin (ATG), cyclosporine A and eltrombopag. Bone marrow transplantation is contraindicated in pregnancy as immunosuppressants are teratogenic to the foetus.

CASE REPORT
Mrs L, 27 years, second gravida with previous normal delivery came at 29 weeks of pregnancy with history of acute onset headache, fever and cough. Her primary gynaecologist had found pancytopenia and had referred her to our haemato-oncology team at Fortis hospital, Bangalore. At admission, her haemoglobin was 4.2 g/dl, her total leucocyte count (TLC) was 2.2 thou/µL (neutrophils 58%) and platelet count 16 thou/µl.

She was hemodynamically stable, supported with intravenous antibiotics piperacillin and tazobactam combination for her fever, packed red blood cells and platelet transfusions. Her iron profile was normal except for marginal raise in ferritin levels. Her LDH, folic acid, vitamin B 12 levels were normal. ANA profile was normal. Bone marrow aspiration showed hypocellular marrow with decreased megakaryocytes. Bone marrow biopsy was consistent with aplastic anaemia.

Her blood investigations in the first trimester were normal with haemoglobin of 11.6 g/dl, platelet count 202 thou/µL, TLC of 6.1 thou/µl (neutrophils 74%). Throughout this pregnancy, besides iron and calcium she did not take any medicines and did not give any history of infections.
Her growth scan at 28th week showed a low-lying placenta with growth corresponding to the gestational age.

A multidisciplinary approach was adopted involving the haematologist, obstetrician, paediatrician, interventional radiologist and a joint counselling was done with the couple. A diagnosis of aplastic anaemia with pregnancy was done and the management plan was discussed with the couple. Bone marrow transplantation with response rates of 70% was discussed. In view of her ongoing pregnancy, this option could be resorted to only if the pregnancy was terminated due to teratogenic effects of immunosuppressant on the foetus. With a 29-week pregnancy and a low-lying placenta, terminating the pregnancy involved risks with preterm caesarean section and bleeding in view of low platelet count.

With the ongoing pregnancy, the treatment options which were now available as per literature review involved (a) Only cyclosporine alone with a success rate of 20%, (b) Anti-thymocyte globulin (ATG) and cyclosporine with a success rate of 66%, (c) Combination of cyclosporine and ATG and eltrombopag- A thrombopoietin agonist with a success rate of 80%.2 In view of her platelet refractoriness a combination of cyclosporine and eltrombopag was offered. However, there was no data on this combination therapy. The couple chose the latter option and she was discharged from the hospital with a haemoglobin of 8.8 g/dl, platelet count of 14 thou/µl and TLC was 5.4 thou/µl, (neutrophils 75%). Her high vaginal swab and urine cultures were negative, antibiotics were stopped.

Treatment was initiated with Cyclosporine 100 mg BD and eltrombopag 150 mg OD. After this, her packed red cells requirements reduced, but she continued to require repeated platelet transfusions. Her liver function tests remained normal. She followed up on outpatient basis with the haematology and obstetric teams and reached 37th week. Her obstetric scan at 37 weeks showed adequate for gestational age foetus with placenta being well away from the OS.

The couple were counselled regarding induction of labour with the involvement of the multidisciplinary team. Her admission blood profile was haemoglobin 7.4 g/dl, platelet count 28 thou/µl and TLC was 5.56 thou/µl (neutrophils 66%). She received a unit of single donor platelets (SDP) and a unit of packed red blood cells (PRBC) one hour prior to induction of labour and one SDP transfusion three hours later. Labour was induced with Propess (sustained release pessary of dinoprostone). She received intravenous piperacillin and tazobactam for antibiotic prophylaxis and delivered spontaneously a live female baby weighing 2.4 kg. Her induction delivery interval was around 7.5 hours. Her labour and postpartum period remained uneventful with no postpartum haemorrhage or infections. Still, we transfused one unit of SDP and one PRBC post-delivery. She remained afebrile and was discharged in a stable condition with haemoglobin of 8.2 g/dl, platelet count 21 thou/µl and total count of 6.2 thou/µl (neutrophils 62%). She was advised to continue eltrombopag and cyclosporine. The newborn blood counts on day three were normal. She did not resort to breast feeding as there were not much data on cyclosporine and eltrombopag. The latter was stopped after four months and cyclosporine was eventually tapered to 25 mg. Now after 14 months post-delivery she is on cyclosporine 25 mg alternate days and with haemoglobin 10.9 g/dl, TLC 4.5 thou/µl and platelet count 135 thou/µl (neutrophils 77%).

**DISCUSSION**

Aplastic anaemia can first present during pregnancy. Spontaneous remission is observed in some women after pregnancy.3,5 An imbalance between placental lactogen, erythropoietin and oestrogen are hypothesised as a cause for aplastic anaemia in pregnancy. The mainstay in the management of pregnancy with aplastic anaemia is supportive care with a multidisciplinary approach involving obstetrician and haematologist.4

In our case, the booking blood tests had revealed normal haemoglobin and leucocyte count. Her symptoms of anaemia got the trigger for performing a bone marrow biopsy. Her initial blood picture had all the components of severe aplastic anaemia with low haemoglobin, platelet and leucocyte count. Her leucocyte count improved within a week. Literature review suggests that neutrophil counts increase during normal pregnancy and that thrombocytopenia is one of the risk factors for complications in pregnancy with aplastic anaemia.5 Platelet transfusions were given daily as her counts hovered around 14, thou/µl (Figure 1). The platelet refractoriness and the high risk of alloimmunisation made it difficult to manage initially. At this stage, she was around 30 weeks, and in the mother’s interest, even if we had considered delivering her with severe aplastic

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**Figure 1:** The trend of platelet count, haemoglobin and absolute neutrophil count through the pregnancy.
anaemia, it would not have been prudent with a placenta lurking close to the internal OS.

**Table 1: Protocol of management of aplastic anaemia during labour**

| Preparation |
|-------------|
| • MD廷 involving anaesthesiologist, gynaecologist, haematologist, blood banking charge, intensivist, intervention radiologist, neonatologist. |
| • Arrangement of donors for platelet and PRBC |
| • Adequate blood stock in blood bank |
| • Arrangement of Novo 7 vials in pharmacy. |
| • Arrange isolation bed in MICU. |

**Role of anaesthetist**

- No lumbar puncture/neuraxial technique for anaesthesia oralgesia.
- No NSAIDS for pain control. Opioids, Nitric oxide acceptable.

**Patient preparation**

- Type of delivery: vaginal preferable.
- Blood thresholds: platelet count >20000 for vaginal delivery, >50000 for LSCS. HB >8 GM/DL. If absolute neutrophil count is 5000 Prophylactic GCSE. Start prophylactic antibiotic Inj. Piptaz 4.5 gm 6 hours before induction/pre-op. (As per our hospital protocol).
- Pre-op/ before induction: 1 SDP transfusion/6 RDP transfusion 1-2 hour before.
- Intra OP 1SDP/6RDP transfusion. 1-unit PRBC transfusion.
- If bleeding continues factor VII (Novo 7) 90 mg/kg slow IV push over 2-5 min.
- Repeat dose of Novo 7 after 6 hours if haemostasis is not achieved.
- Post op/post-delivery 1 SDP/ 6 RDP transfusion.
- Monitor platelet 12 hourly, transfuse 6 units RDP/1-unit -SDP to maintain platelet count >20000 for 5 days post OP.
- Treat fever aggressively as per febrile neutropenia protocol.

Induction of labour was planned for logistic reasons to maintain a good platelet count and haemoglobin. As her leucocyte counts were normalised, induction of labour with vaginal insertion with Propess was considered to be safe. Propess is a sustained release vaginal delivery system of dinoprostone 10mg.

Considering her platelet count and haemoglobin improved after delivery, pregnancy may have been a triggering event.

**CONCLUSION**

A team approach and robust support from the family played a key role in successfully managing our case. The leucocyte count recovered early and hence infections did not pose a threat. This patient seemed to be very refractory to platelet transfusion and needed several platelet transfusions initially.

Eltrombopag probably is a wonder drug to improve outcome in pregnancy with aplastic anaemia but safety in pregnancy needs to be ascertained.

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