Case Report

HELLP syndrome: An experience of treating with plasma exchange
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Abstract:
In the spectrum of patient with severe pre-eclampsia, there is a potentially lethal complication called HELLP syndrome (haemolysis, elevated liver enzymes and low platelet count). The pathogenesis of HELLP syndrome is unclear. If it is a form of severe preeclampsia, it likely originates from aberrant placental development and function. As an independent entity, it has been attributed to abnormal placentation, similar to preeclampsia, but with greater hepatic inflammation and greater activation of the coagulation system than in preeclampsia. HELLP develops in approximately 0.5% to 0.9% of all pregnancies in 10% to 20% of women with severe preeclampsia. Maternal mortality rate in HELLP syndrome is variable (1-23%) due to severity of disease, delayed diagnosis and presence of multi-organ involvement. So the recognition of HELLP syndrome and an aggressive multidisciplinary approach are required for the improvement of maternal foetal prognosis.

Key Words: HELLP, Preeclampsia, Therapeutic Plasma Exchange.

Introduction:
HELLP syndrome and severe preeclampsia are probably part of the same disease spectrum. A precise definition of HELLP is necessary for research purposes and for predicting maternal complications as well as for deciding treatment modality.

As per Tennessee Criteria, we require the presence of all of the following to diagnose HELLP:9

- Microangiopathic hemolytic anemia with characteristic schistocytes (also called helmet cells) on blood smear. Other signs suggestive of hemolysis include an elevated LDH or indirect bilirubin and a low serum haptoglobin concentration (<25 mg/dL).
- Platelet count <100,000 cells/µL.
- Serum LDH >600 IU/L or total bilirubin >1.2 mg/dL.
- Serum AST >70 IU/L. Some investigators obtain ALT levels instead of, or in addition to, AST levels. An advantage of the AST is that it is a single test that reflects both hepatocellular necrosis and red cell hemolysis.9

A less established criteria known as Mississippi Criteria; for prognostication of patient also exists which divides the patients into three groups based on lowest observed perinatal platelet count. Class 1 platelet nadir <50,000/cmm, Class 2 >50,000 - 100,000/cmm and Class 3 >100,000 - 150,000/cmm. This criteria along with multiorgan involvement may serve as a deciding factor for initiating plasma exchange.

Here we report a case diagnose to have HELLP syndrome and its successful management with plasma exchange.

Case report:
A 32 years, young women at her 38 weeks of gestation was admitted to ICU, SHL with epigastric pain, blood pressure 160/120 mm of Hg and proteinuria. She underwent emergency LSCS at the same day due to severe preeclampsia and foetal distress.

After surgery, she was conscious and alert without any neurological deficit. Her blood pressure remained elevated up to 160/120 mm of hg and required i/v antihypertensive. She was maintaining SPO2 94% with 4lt/min O2. Abdomen was distended and diffusely tender. There is bleeding from venepuncture sites. Her initial blood counts in post operative period was within normal range. On 2nd post operative period she developed severe thrombocytopenia (platelet 22X103/cmm with hemoglobin of 9.8 gm/dl and white blood count 22X103/cmm. Her coagulation profile was normal. Peripheral smear showed significant numbers of schistocytes suggestive of microangiopathic hemolytic anemia(MAHA). Elevated ALT of 1104 IU and LDH of 12695 U supported the diagnosis of MAHA. Culture of urine, sputum and blood that were sent for septic screening before starting antibiotic, turned out to be negative later on. Hepatitis viral markers (HAV, HBV, HCV & HEV) were found negative also. Transabdominal USG revealed mild intraabdominal collection, with no organ injury. To rule out
autoimmune disorder, antinuclear antibody, antineutophilic cytoplasmic antibodies were tested and found to be negative.

Her clinical and biochemical condition deteriorated in 3rd post operative day. Patient became disoriented and urine output decreased to 100 ml in 24 hours. Repeat hematological investigation revealed persistent thrombocytopenia and Hb dropped to 7 gm/dl and LDH level increased to 13,500 U. Her creatinine was also rising. A provisional diagnosis of post partum acute renal failure due to HELLP syndrome was made. Hence, in view of the clinical deterioration an emergency therapeutic plasma exchange (TPE) was carried out after 48 hours of presentation. Four session of plasma exchange was done on a daily basis. As she became anuric & renal function deteriorated hemolysis was initiated. Supportive treatment with PRBC transfusion was also given when deemed necessary.

After 4 days of TPE, clinical and biochemical condition was improved and she was shifted from ICU to regular ward. She started having urine output after 7 days and that gradually increased and HD could be stopped at day 10 of presentation. She was discharged from hospital at 15 days of her hospital course with near normal renal function and without any evidence of ongoing hemolysis and normal platelet count.

Discussion:
Many aspect of HELLP syndrome are still controversial, such as diagnosis, classification and management.10,11 During the post partum period, the majority of cases will show resolution of the disease within 72 hours of delivery; however, some patient , especially those with the persistent form of the disease, will not improve or will even get worse. Some authors have suggested that such cases might get benefit from plasma exchange.12 The exact mechanism of the effect of plasma exchange in HELLP syndrome is not known, but in general, plasma exchange removes plasma along with inflammatory marker and autoantibodies from patient and substitutes new elements from donor plasma. There is thin line of demarcation which differentiates HELLP syndrome from other thrombocytopenia associated multiple organ failure like TTP/HUS. In this case, patient was diagnosed as a case of HELLP syndrome. As patient's clinical and biochemical condition was deteriorating in post partum period and diagnosed as class-I HELLP syndrome and had developed multiorgan failure, TPE was initiated and patient improved rapidly and her renal recovery took also significantly less time.

Although large scale controlled trials are not available till date and further study is needed; Therapeutic Plasma Exchange could be an option in patients refractory to standard management or who have Class I HELLP or develops multiorgan failure.

References: