Management of concurrent severe preeclampsia and thrombotic thrombocytopenic purpura (TTP)

Abstract
Thrombotic thrombocytopenic purpura (TTP) is a devastating disease involving the micro circulating (capillary wall damage) of multiple organs and could be seriously fatal if not diagnosed and treated promptly.1 TTP is a rare complication of pregnancy with a poor prognosis and high fetal mortality, especially when it occurs during the first trimester.2 But most of TTP cases occur in the late trimester of pregnancy. Untreated TTP is associated with more than 90% mortality rate.3 Plasma replacement, even in pregnancy, remains as the corner stone of prevention and treatment of TTP, especially those with autoantibody- driven ADAMTS-13-deficient TTP.

Keywords: complication, treatment, proteinuria, hypermagnesaemia, epigastric

Case report
A 24-year-old parturient G2 P1 A0 at the gestational age of 38 weeks, presented at emergency department. She had a history of proteinuria (3+) eight months prior to admission in our hospital. Because of severe high blood pressure 220/160mmHg, proteinuria (3+), high serum creatinine (cr.=1.7), mild loss of consciousness and she was admitted in intensive care unit. After some medical treatment (magnesium sulfate and hydralazine) and partially stabilization of her blood pressure, urgent pregnancy termination via lower segment caesarean section (LSCS) was performed. On admission in ICU (the first day) she had no headache and blurred vision, but complained of epigastric pain and nausea. Blood sugar was normal vital signs: BP:220/160mmHg, PR-90/min, RR:18/min, T:37.8 C., spo2>92% (by room air).

Coagulation tests (PT, PTT and INR) were all within normal limits on the admission time and all over the course of hospitalization. Urine analysis showed RBC=3–4 and proteinuria (3+). The cell blood count showed result: WBC:13000, Hb:11.5, Hct:35, MCV:89.75, PLT:181000, RDW:13.2.

Blood pressure was controlled by hydralazine 5mg, IV/repeated doses and methyldopa 250mg/q4h PO. Labetalol was not with reach at that time. Mgso4 was given promptly as a 4gr loading dose intra venous, followed by a maintenance intravenous infusion of 1 to 2gr/h by controlled infusion pump. It lasted just for 24 hours and at the rest of the course of treatment we had to discontinue Mgso4 infusion because of severe hypermagnesaemia. On the day two of ICU admission (one day after caesarean section) the patient had hemoptysis, thrombocytopenia (PLT:56000) hyperkalemia (k:6), high lactate dehydrogenase (LDH:3125), Hyperbilirubinemia (Bill Total:3.3;Bill Direct:1.3), peripheral blood smear showed anisocytosis, Burr cell, Roulex formation mild increase of liver function test (AST=47, ALT=80), negative blood cultures after 48 hours, decrease of HB up to 8.5mg%, rise of creatinine up to 3 and more, mild loss of consciousness; all were in favor of HELLP syndrome or TTP concomitant with severe preeclampsia. According to hematologic consultation, our patient was supposed to undergo plasma exchange after verifying the existence of schistocytes in peripheral blood smear. Hence, plasma exchange was requested for the patient straight after PBS examination. D-Dimer>10000, FDP>20, Serum Fibrinogen:165.

Unfortunately 5 units of platelets and 10 units of cryoprecipitate were transfused to the patient soon after C/S in recovery room. Furthermore thrombocytopenia started to become worse. On days three and four of hospitalization we were facing additional problems, making the current condition worse, these were oliguria, profound thrombocytopenia PLT. Count:25000, Fibrinogen:270, FDP=20, massive bloody secretions from the drain inserted in caesarean incision, refractory metabolic acidosis(PH<7.3), Cr>3 and all findings in favor of acute kidney injury(AKI)-both in laboratory tests and ultrasonographic examination- and also gross hematuria, hypermagnesaemia and severe ileus resulted in abdominal distension and respiratory distress(Ileus treatment was performed immediately). Liver function tests (ALT&AST) were increased up to 94. In this new condition a shaldon catheter was placed in right subclavian vein and the patient underwent hemodialysis. On day three we had a report of slight amount of schistocyte and (+1) on the next day, hence, plasma exchange was requested for her on a daily basis. On the other side, she was on daily hemodialysis without heparin, transfusion of whole blood, FFP, Cryoprecipitate, antihyptertensive (consisting of calcium channel blockers i.e. diltiazem, α blockers i.e. prazocin), anticonvulsives, antibiotics including ceftriaxone and clindamycin, steroid therapy, intra vascular fluid therapy. All of these were on the basis of the progress of patient’s condition. On day 6patient developed polyuria urine output: 250-300c/c/h. On the Other side we had a rise in platelet count up to 80000 and never decreased from that time. Bleeding from the drain lessened obviously. On day 7patient’s consciousness was complete with oriented voluntary movement of limbs. At that time we had no schistocytes on PBS, decreasing of LDH less than 800(LDH<800) and no bleeding tendency, therefore plasma exchange was discontinued.
On day 10, after keeping tight control on fluid and electrolyte balance during polyuric phase of AKI, patient recovered her consciousness and showed a decreased in serum creatinine level and correction of acidosis. Hence, there was no more need to hemodialysis. On day 13 the patient began to breast feed her child. Plasma exchange discontinued. On day 17 there was no need to hemodialysis and the patient was good enough to be discharged from the hospital.

Additional diagnostic procedures and tests

Since the patient was too ill to be transferred to CT-scan department all diagnostic procedures were performed portable on the bedside. We had the chance of having definite serial ultrasonographic checks of abdominal and pelvic cavities indicating that we could be sure of not having any kind of abnormal internal bleeding. This fact helped our obstetrician not to perform second revision of the site of operation. Laboratory findings including: ANA, FANA, Anti dsDNA, APA, C3, C4, CH50, HBS Ag, Anti HCV Ab, Anti HIV, lupus anticoagulant, anticardiolipin and serum albumin were not abnormal. ADAMTS 13 Ag 5.22 high ADAMTS 13 auto Ab 14.4 borderline verifying in blood sample. Complications due to invasive procedures at the time of patient’s presentation to hospital, she was suffering from mild to moderate respiratory distress investigation by chest x-ray and chest sonography indicated moderate bilateral pleural effusion. In addition after replacement of shaldon catheter on the day of admission, we had right sided hemopnuemothorax leading to insertion of chest tube, on day 9, chest tube discharge was less than 100cc/24h and on day 12 we removed it. Biopsy sample taken from kidney 3 months after hospitalization. The glomerulus shows segmental sclerosis with syneciae to overlying, Bowman’s capsule (methenamine silver, x400) (Figure 1, Figure 2 & Figure 3).

Discussion

There are several fatal conditions complicating preeclampsia and should be under consideration seriously, consisting of systemic lupus erythematous (SLE), anti-phospholipid syndrome (APS), acute fatty liver during pregnancy (AFLP), thrombotic microangiopathies (TMA) appearing as thrombotic thrombocytopenic purpura (TTP) or hemolytic uremic syndrome (HUS), HELLP syndrome and sever sepsis (5-7). Among these imitators of preeclampsia the differential between TMA and HELLP syndrome is really confusing and play an important role in saving the mother and her future child’s lives. According to some studies the definitive marker for TTP is undetectable blood ADAMTS 13 activity.5–10 In general terms, some complications during pregnancy including hemolysis, concurrence of arterial hypertension and proteinuria, renal failure, elevated liver enzymes, thrombocytopenia and in some cases signs of visual, neurological, pancreatic or pulmonary disturbances lead to misdiagnosis in severe preeclampsia Complicated with HELLP syndrome or with thrombotic microangiopathies (Table 1).11

Table 1 Clinical and Laboratory Features of TTP, HUS, and HELLP

<table>
<thead>
<tr>
<th>Clinical and laboratory features</th>
<th>TTP</th>
<th>HUS</th>
<th>HELLP</th>
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<tr>
<td>Neurologic sxs</td>
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<tr>
<td>Fever</td>
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<tr>
<td>Hypertension</td>
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<td>Renal dysfunction</td>
<td>+/-</td>
<td>+++</td>
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<tr>
<td>Skin lesions-purpura</td>
<td>+</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Platelets</td>
<td>↓↓</td>
<td>↓↓</td>
<td>↓</td>
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<tr>
<td>Prothrombin time (PT)/activated partial thromboplastin time (aPTT)</td>
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<td>⇔</td>
<td>↑ or ⇔</td>
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<td>Fibrinogen</td>
<td>⇔</td>
<td>⇔</td>
<td>⇔ or ⇔</td>
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<tr>
<td>BUN/creatinine</td>
<td>↑</td>
<td>↑↑</td>
<td>↑ or ⇔</td>
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<tr>
<td>Aspartate aminotransferase (AST)/alanine aminotransferase (ALT)</td>
<td>⇔</td>
<td>⇔</td>
<td>↑</td>
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<td>Lactate dehydrogenase (LDH)</td>
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Figure 1 The glomerulus shows segmental sclerosis with syneciae to overlying, Bowman’s capsule.

Figure 2 The glomerulus shows segmental sclerosis with syneciae to overlying, Bowman’s capsule.

Figure 3 The glomerulus shows segmental sclerosis with syneciae to overlying, Bowman’s capsule.
In any case all appropriate treatments of preeclampsia (antihypertensive-magnesium sulfate) and other initiators of it should be commenced without delay. Since TTP has got a high mortality rate if not treated immediately, once the diagnosis of it is certain or even highly suspected, plasma exchange should be considered as the matter of great urgency, often in conjunction with steroid, antibiotics, when there are clues of severe sepsis. Platelet infusion should be avoided, even in cases of profound thrombocytopenia, as it could worsen the situation and lead to sever cardiovascular events in TTP. Many papers have dealt with this question that, whether sever HELLP syndrome particularly in the postpartum period is a form of TMA or not? R,5,6,15-19 Having detailed knowledge of natural history of each disease process and the considerable overlap of them, which make them seem to be “disease syndrome” rather than single disease, will certainly help us to distinguish these related peculiar conditions. Considering to all mentioned above we should bear in mind not to misdiagnose TTP as HELLP syndrome.6,15 More over regarding to the risks of large volume plasma, plasma exchange should be requested just when there is highly suspected evidence of TTP/HUS.19 In addition to all previous concerns, the financial cost of plasma exchange constitutes a great part of its considerations.8-12 Among all similarities between HELLP syndrome and TMA, some are in favor of TMA and the others were indicating HELLP syndrome existence.

One helpful hint is that hemolysis is at foreground In TTP, HUS, leading to red-colored urine or/and resulting the need for red blood cells transfusion. On the other side, in HELLP syndrome, the intensity of hemolysis is far less than TMA. It is emphasized by Weinstein himself, that hemolysis was absent in about 30% of patients.20 One another point in differentiation of these two conditions is profound thrombocytopenia in TMA, as low as 10-20*10⁹L⁻¹, comparing to HELLP syndrome in which platelet count often remains more than 50*10⁹L⁻¹, the lower amounts happen in about 15% of all patients.8 There are five groups of signs which are called the classic pentad of TMA, emphasizing that TTP is a mechanical hemolytic anemia. These are thrombocytopenia, microangiopathic hemolytic anemia (coombs negative) ruling out autoimmune hemolytic anemia-fever, neurologic disturbances and renal involvement which are fully presented in only a minority of patients. Concurrent existence of hemolytic anemia and thrombocytopenia is indicative of TTP. Three other less common classical manifestations are: renal failure, central nervous involvement and fever.13,21-23 We should keep in mind that in new cases of severe hemolysis associated with profound thrombocytopenia, laboratory data should be obtained to rule out TMA.3,13,14 Recently one proposed laboratory finding for TTP is a lactate dehydrogenase (LDH) /aspartate aminotransferase (AST) ratio above 22.12,19 In the past pathological criteria were mandatory to define the disease, consisting of glomerular capillary endotheliosis associated with micro vascular platelet thrombi, fibrin formation in and around glomeruli, widespread arterial thrombosis, sub endothelial clear deposits in glomerular capillaries.3,13,14,22,23 Since a vast majority of patients show profound thrombocytopenia and critically ill there is a limited access to histological studies. Findings in favor of TTP are: micro vascular thrombi of Von Willebrand factor (VWF) and fibrin in kidney, skin or bone marrow, while fibrin thrombi alone indicates HELLP syndrome more than the other possibilities. Current trend is towards diagnosis of TTP by measuring a biological parameter (ADAMTS 13 activity), if less than 10% (in case of TMA highly imitating HELLP syndrome) is known as an index for TTP.13,14 The activity of this enzyme is low in HELLP syndrome too, but not to the same extent as in TTP. In some papers it is reported to be 31% (range 12-43%) in HELLP syndrome, compared with 71% at the end of normal pregnancy and less than 10% in TTP.24,25

It is advisable to make treatment decision with the help of this index.24 Some other useful factors including severity of liver disturbances in comparison to hemolysis and dramatic rise in liver function tests are highly suggesting of HELLP syndrome more than the other diagnosis. On the other side data indicating mild liver involvement very high level of LDH, elevated LDH/AST ratio and gross hematuria support the existence of TTP.24 Some researchers have said that in cases of ADAMTS13 activity within the range of usually seen at the end of pregnancy,29,30 in other word mild decrease in ADAMTS13 activity allows the rule out TTP, and post-partum severe HELLP syndrome can be proposed. Hence this situation can be resolved without the need for plasma exchange, but we cannot ignore the possibility that plasma exchange may be of some benefit in post-partum severe HELLP syndrome after all.29-31 Yet, the question about plasma exchange utility in these situations remains unanswered. Very recently, there is report of nine cases successfully underwent plasmapheresis by Owens et al., those were defined as “post-partum thrombotic-angiopatihic syndrome.32”

ADAMTS13 testing

TTP is a rare disease (5-10cases per million persons per year) characterized by the massive formation of platelet rich-thrombi in the microcirculation of multiple organs.30,34 In plasma of patients with recurrent TTP because of the deficiency of a cleaving protease ADAMTS13 as being responsible for the presence of ultra large (UL) multimers of Von Willebrand factor in endothelial cells, platelets and plasma.39 After stimulation of endothelial cells UL forms of VWF present in the endothelium but not in plasma in normal conditions, promote intravascular aggregation of platelets and the consequent micro vascular and mechanical hemolysis resulting in high fluid shear stress especially in the microcirculation.39 A kind of protease responsible for regulating the multimeric structure VWF, which was identified in 2001 by Zhang et al.17 In a retrospective cohort study of patient with TTP the VWF cleaving protease was found deficient.30,31 The new member of the ADAMTS (a disintegrin and metaloprotease with thrombospondin 1 repeats) family of metaloprotease was called ADAMTS13.34

Classification of TTP

TTP has two main forms congenital and acquired. The first one is due to mutations in the ADAMTS 13 gene; it is rare (1:1000000) autosomal recessive and most of the time manifests at birth or during childhood.40,41 The second one which is acquired is found in two types: immune-mediated forms, due to auto antibodies against ADAMTS 13.42,43 and the other one is caused by massive endothelial stimulation with consequent release of UL VWF multimers in extremely large amount exceeding the system’s ability to degrade them, though the levels of ADAMTS 13 level can be normal or mildly reduced.44 The most commonly conditions present in the immune-mediated forms, often associated with sever ADAMTS 13 deficiency (levels less than 10% of normal) are pregnancy, infections, autoimmune disease and the use of drugs consisting of ticlopidine and clopidogrel.

On the other side the most common conditions associated with the type of TTP presenting with normal or mildly reduced levels of ADAMTS 13 (greater than 10%) are metastatic tumors, organ
transplantation and the use of drugs such as cyclosporine, mitomycin and a-interferon. According to severe recent studies those kind of TTP associated with severe deficiency of ADAMTS 13 activity less mortality rate than those with detectable ADAMTS 13 activity. By contrast, Coppo et al. reported much high mortality rate in patients with severe ADAMTS 13 deficiency in comparison to those with non-severe deficiency. Higher mortality rate of patients with detectable ADAMTS 13 activity could be related to the lethal fault of their underlying diseases. Along with aforementioned studies, many researchers have been carried out to define. Define the strong relation between inhibitory anti ADAMTS 13 testing and mortality rate. Overall, we come to the conclusion that existence of anti ADAMTS 13 is associated with a worse prognosis. Moreover there are some reports of a positive correlation between high inhibitor titers and severity of clinical manifestations, treatment refractoriness and the rate of death.

Conclusion

The presence of schistocytes (fragmented red blood cells) on the peripheral blood smear suggest red blood cell injury from damaged endothelium and is a characteristic feature of microangiopathic hemolytic anemia. These are coombs-negative intravascular hemolytic anemia including TTP and HUS. In these two situation coagulation factor assays (prothrombin or partial thromboplastin time, fibrinogen, D-dimer and soluble fibrin monomer) are not abnormal, at least not in the initial stages. Opposite to it there is often an acute-onset profound thrombocytopenia (plt. Count, <20*109/L), especially in the presence of renal, CNS involvements which should keep in mind the diagnosis of TTP, HUS and if confirmed immediate specific therapy (plasma exchange) must be performed. In this article we presented a case of acute TTP secondary to acquired auto antibodies complicated by severe preeclampsia treated by plasma replacement of (40-50cc/kg plasma) were removed at each procedure and substituted with FFP, to replace ADAMTS 13 and clotting factors, concomitant with methylprednisolone and daily hemodialysis. Signs and symptoms of preeclampsia and TTP were treated by degrees to save the parturient and child lives. On admission time, things were too confusing to make a straight forward decision but as a matter of fact some clues consisting of worsening of thrombocytopenia due to platelet infusion on the day of C/S, detecting fragmented RBCs on PBS, dramatic rise of LDH but mild rise of LFT, coombs test; ADAMTS13 assays (anti ADAMTS13 inhibitor more reliable than the other tests), mild abnormality in liver function tests, coagulation factors assays range within normal limits (at least at the early stage of TTP); can help clinicians to distinguish TTP from HELLP syndrome. However if there is any doubt about TTP existence we should lose no time commencement of plasma exchange. Furthermore some physicians believe in adjuvant therapy by steroids or immunosuppressive agents. Still lots of encouraging studies are in process to maximize the quality of ADAMTS13 assays to define TTP.

Acknowledgements


Figure 4 Platelets level.

Figure 5 Creatinine level.

Condensation

In all cases of severe preeclampsia, severe HELLP syndrome and TTP, we should not hesitate for the appropriate treatment in all aspects of the disorders including preeclampsia (magnesium sulfate, antihypertensive- anticonvulsant) and the disease or in other word the micro angiopathic syndrome concomitant with it. Some findings consisting of, dramatic rise in LDH, profound thrombocytopenia not responding to platelet infusion. Fragmented red blood cells on peripheral blood smears, ADAMTS13 assays (anti ADAMTS13 inhibitor more reliable than the other tests), mild abnormality in liver function tests, coagulation factors assays range within normal limits (at least at the early stage of TTP); can help clinicians to distinguish TTP from HELLP syndrome. However if there is any doubt about TTP existence we should lose no time commencement of plasma exchange. Furthermore some physicians believe in adjuvant therapy by steroids or immunosuppressive agents. Still lots of encouraging studies are in process to maximize the quality of ADAMTS13 assays to define TTP.
Conflict of interest

The author declares no conflict of interest.

References

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Information about the content of the image is not provided. However, based on the metadata and the structure of the document, it appears to be a scholarly article discussing the management of concurrent severe preeclampsia and thrombotic thrombocytopenic purpura (TTP). The content includes references to various studies and clinical observations related to the diagnosis and treatment of these conditions. The article likely discusses the complexities of managing a patient with both conditions, possibly highlighting case studies, treatment strategies, and outcomes from clinical trials or case series. The references cited in the text suggest a comprehensive approach to understanding the pathophysiology, laboratory markers, and therapeutic interventions for these conditions.