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The Use of Plasma Exchange in a Very Early-onset and Life Threatening, Hemolysis, Elevated Liver Enzymes, and Low Platelet (HELLP) Syndrome: A Case Report


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Abstract

Background: HELLP syndrome is a life threatening pregnancy and early postpartum complication. Very early presentation (before the 21st pregnancy week) is rare and represents an extremely difficult situation for patients and physicians. Supportive therapy (magnesium sulfate, antihypertensive drugs and corticosteroids) may be useful to prolong pregnancy; till now the removing of placenta is the only effective therapeutic option. Plasmapheresis may represent a new and efficacious therapeutic option.

Case description: The article reports on a case of very-early onset HELLP Syndrome at the 18th (17+5) week of gestation. It was a challenging clinical and therapeutic case. Since the fetus did not show any signs of growth retardation or pathological Doppler findings, indicating a good fetal prognosis, we used plasmapheresis as an ultima ratio to prolong pregnancy. With plasmapheresis, the pregnancy was prolonged for 20 days. Unfortunately the deterioration of the clinical situation required delivery in the 21st (20+4) gestational week.

Conclusion: Plasmapheresis allows prolongation of pregnancy with early onset, life threatening HELLP-Syndrome.

Keywords: Management of early HELLP syndrome; Plasma exchange; Ultrasound; Imitators of HELLP syndrome

Introduction

The HELLP syndrome occurs in about 0.5 to 0.9% of all pregnancies and in 10 to 20% of cases of severe preeclampsia [1]. In about 70% of cases, the HELLP syndrome develops before delivery [2]. Almost 10% occur before the 27th week of pregnancy [3]. Presentation before the 20th week of pregnancy is a rare condition [4]. Maternal mortality is estimated from 1.1% to 25% [5]. The causes of death are mostly cerebral hemorrhage and disseminated intravascular coagulopathy [6]. Perinatal mortality depends on the gestational week and is estimated between 7.4% and 34% due to prematurity, placental insufficiency and abruptio placentae [7].

The pathogenesis remains multifactorial and is not completely understood. Immunological maladaptation and aberrant invading trophoblast are involved in the development of both early-onset preeclampsia and HELLP syndrome. Anti-angiogenetic factors are released from the placenta into maternal circulation inducing the symptoms: sFlt-1 (soluble fms-like tyrosine kinase 1 or soluble vascular endothelial growth factor -VEGF- receptor 1) levels are similar in preeclampsia and HELLP syndrome, other mediators like Fas Ligand and endoglin may be responsible for the greater hepatic inflammation and activation of the coagulation system in HELLP syndrome cases [8]. Together with the increased levels of sFlt-1, there are reduced levels of VEGF and placental growth factor (PIGF), resulting in endothelial dysfunction [9].

Weinstein first described this syndrome in 1982 [10]: “H” represents hemolysis, “El” elevated liver enzymes and “LP” low platelet count. There exists also an incomplete form, with just one or two elements of the triad [1].

Clinical management depends on the gestational age. Delivery is indicated when the syndrome develops after the 34th gestational week. Delivery after 48 hours of evaluation and treatment with corticosteroids appears reasonable between the 27th and 34th week of gestation. If the maternal or fetal conditions deteriorate, delivery should be performed as soon as possible. If the laboratory results or the clinical condition deteriorate, postpartum plasma exchange is reported to improve maternal outcome of severe HELLP syndrome [11].

Case Report

We present the case of a 28-years-old Caucasian primigravida at 17 weeks and 5 days of pregnancy who was admitted to a local hospital with complaints of nausea, pain in the chest and in the right upper abdominal quadrant. An electrocardiography and echocardiography were performed and these were not pathological. She was then referred to our tertiary referral centre because her laboratory studies revealed probably a very early HELLP Syndrome: platelet count 40 nl (normal range 180-380), Hb 10 g/dl (12-15.2), ASAT 141 U/l(<35), ALAT 254 U/l(<35), LDH 376 U/l (120-247), Haptoglobin 0.07 g/dl (0.4-2.4). At admission, the blood pressure was normal 120/70 mmHg under the...
administration of methyl dopa 250 mg twice daily since 1 week. She
denied a history of chronic hypertension, but she had one hypertensive
episode during a recovery in a psychiatric ward because of depression
one year before the pregnancy. She revealed the use of cannabis up
until in the early pregnancy. She denied the use of cannabis or other
illicit substances after knowing she was pregnant. She smoked 4
cigarettes a day. The physical examination revealed no jaundice and no
evidence of purpura or petechia, no evidence of neurologic
abnormalities or edema, and she was able to converse at her baseline
mental status. Her deep tendon reflexes were normal. Fetal cardiac
activity was appreciated by ultrasound examination, the Doppler
studies of the uterine artery showed pathological values: PI left 2.53
and right 3.50 with notching on both sides.

The administration of methyl dopa was continued and an
intravenous seizure prophylaxis with magnesium sulfate 1 g/h as well
as steroid administration was started (10 mg Dexamethasone 3 times a
day for 2 days).

Considering the early and unusual presentation, a comprehensive
process to exclude differential diagnoses was started. The case was
interdisciplinary discussed with the colleagues from the departments
of internal medicine, nephrology and gastroenterology. An ultrasound
examination of the upper abdomen did not demonstrate any
pathological relevance in this region. Normal activity of ADAMTS 13
was registered (80%, normal range 50-110%), excluding thrombotic
thrombocytopenic purpura. Renal function was normal (Creatinine 0.6
mg/dl, normal range 0.6-1.1). There was no icterus, no neurological
deficits, normal bilirubin levels (0.8 mg/dl, normal range 0.3-1.2),
normal level of fibrinogen (350 mg/dl, normal range 150-400), slightly
reduced ATIII activity (70%, normal range 80-120) and other
coagulation parameters were within normal range. Autoantibodies were
in normal range: ANCA IFT <1:10 (<1:10), HepIFT 1:80 (<1:80),
Anticardilipin IgM 1.3 and IgG 1.4 (<7 und <1:10). sFlt-1 at admission
was high, 25292 pg/ml.

After the therapy with corticosteroids, the situation was stabilized
and laboratory values were nearly normal (Thrombocytes 170/ni,
ASAT 39 U/l, ALAT 136 U/l, LDH 376 U/l). The sFlt-1 value drops to
13410 pg/ml and the sFlt-a/PIGF Ratio was 353.2). Unfortunately, 4
days since the admission, the laboratory studies and the clinical
condition deteriorated once again.

The blood pressure became higher with edges of 180/100 mmHg
(Figure 1), hemoglobin fell to 7.8 g/dl, liver enzymes became higher
with ASAT 1141 U/l, ALAT 935 U/l, yGT 124 U/l, LDH 1591 U/l and
thrombocytes 19/nl (Figure 2). The urine examination showed a
significant proteinuria (440 mg/24h). A drug screening detected the
presence of amphetamines in urine. A chromatography blood test
excluded the assumption of such drugs. A forensic evaluation
considered it as a false positive urine screening due to cistitis. A
pheochromocytoma was excluded.

Blood pressure was kept under control with intravenous application of
urapidil (till 20 mg/h), metoprolol 47.5 mg twice daily further to
methyl dopa 250 mg 4 times daily. The patient aimed to prolong the
pregnancy. The fetus showed normal growth curve with normal fetal
Doppler flows (umbilical artery PI 1.40, arteria cerebri media PI 1.31,
peak systolic velocity of 36.9 cm/s) and no signs of any fetal
abnormalities. After consulting the nephrologist and obtaining
informing consent of the patient, plasma exchange was started.

14 sessions of plasma exchange were conducted in total with the
Spectra Optia® and Cobe® Spectra Apheresis System. The estimate of
plasma volume (EPV) was calculated using the formula of Kaplan AA
[12] for EPV. EPV = [0.065 × body weight(kg)] × [1-Hematocrit]. For
anticoagulation acid citrate dextrose-A (ACD-A) was used. A dialysis
catheter was positioned on the right neck. Before starting,
presinolone, clemastine und cimetidine was administered intravenously. Plasma was replaced with 2,5 l of fresh frozen plasma
form a donor.

First, we noted an improvement of the general condition of the
patient and laboratory results (Figure 3). During the plasma-exchange
and 14 days after admission, sFlt-1 concentration in serum drops
further to 10100 pg/ml, but also PIGF also drops (21,7 pg/ml) with an
increased ratio of 466,5. However, since the determination of sFlt/
PLGF-ratio was not clinical routine during the time of therapy, we
failed to determine these values daily during therapy.

The clinical condition of the patient, blood pressure and the
laboratory findings worsened again (Figure 2). She developed
pulmonary edema, dyspnea and the blood pressure was no longer
under control with edges of 200/110.

After extensive interdisciplinary discussions, removing the placenta
was the only therapeutic option left to improve the HELLP syndrome.
Because of the patient’s bad condition, an immediate cesarean section
was performed at 21st week of pregnancy, after 20 days of
hospitalization. The female fetus was delivered which weighted 310 g
with a head circumference of 16.5 cm and a length of 23 cm. She was
assisted by the pediatricians and died 2.5 hours postpartum.
The histological examination of the placenta showed very small terminal
villi without sufficient vasculatization. After the operation, the patient
remained in the intensive care unit for 2 days. The further
postoperative course was uneventful.

After the delivery, the condition of the patient improved rapidly. At
discharge, antihypertensives medication consisted of metoprolol and
amlodipine.

Discussion
This case underlines two important issues in the about management
of the very early HELLP syndrome:

- Exclusion process of imitators of HELLP Syndrome and substance
  abuse
- A new therapeutic option with the plasma exchange

Severe preeclampsia and HELLP syndrome can resemble a number
of different microangiopathic disorders, making it a hard challenge for
the physician to distinguish between them.

The clinical presentation of HELLP syndrome commonly includes
nausea, vomiting, epigastric pain. In 85% of the cases, the signs and
symptoms of preeclampsia such as hypertension, proteinuria,
headache, visual changes and edema are also present.

The symptoms of acute fatty liver in pregnancy (AFLP) are almost
identical with those of HELLP syndrome, but women with AFLP can
rapidly develop liver failure and encephalopathy. Hypertension and
proteinuria are more frequent in HELLP syndrome, but only the
laboratory can help to better distinguish between the two entities: in
AFLP a prolongation of prothrombin (PT) and of activated partial
thromboplastin time (aPTT) is present. Further, ammonia is more
frequently elevated and bilirubin and creatinine are always too high.
Liver biopsy is recommended as the standard procedure to confirm
the diagnosis, but requires an acceptable haemostatic function [13].
Microangiopathic disorders such as thrombotic thrombocytopenic purpura (TPP) and hemolytic uremic syndrome (HUS) are extremely rare in pregnancy (1:100,000).

The classic manifestations of TTP include thrombocytopenia, microangiopathic hemolytic anemia, neurological abnormalities, fever and renal failure with the first three signs present in almost 50-75% of the cases. In TTP the activity of the von Willebrand factor cleaving metalloprotease known as ADAMTS-13 is reduced. The elevated levels of large multimers of the von Willebrand factor induce platelet aggregation and endothelial injuries. Patients with TTP present a strong reduction of ADAMTS-13 activity (<5%), whereas in HELLP syndrome the metalloprotease activity is normal or just slightly reduced.

HUS is an exceptional event during pregnancy and it is mostly seen in children after enteric infection with E. coli that produces Shiga toxin. Typical manifestations are edema, hypertension, bleedings and severe renal failure. This last feature is more severe than in the others microangiopathic disorders [13].

The treatment of HELLP syndrome consists of the delivery of the placenta. Nevertheless, the aim of treatment is prolongation of pregnancy and symptomatic therapy if deterioration or first signs appear after delivery. Hypotensive drugs and seizure prophylaxis with magnesium sulfate represent the cornerstones of the treatment. The effectiveness of dexamethasone is not clearly defined [14]. However, some research demonstrated that postpartum plasma exchange therapy improves treatment outcomes in patients with severe HELLP syndrome [11,15-17]. Patients who do not show improvement of platelet and ASAT levels within the first 24 to 48 hours after delivery and patients with renal or neurological involvement may have benefit from plasma-exchange between 24 and 72 hours postpartum as shown by Simetka et al. [18].

Antepartum plasmapheresis for HELLP Syndrome was first described by Martin et al. seven gravidas between 24 and 30 weeks of gestation with severe preeclampsia were recruited for the treatment from 1984 to 1987. Maternal-fetal deterioration required cesarean delivery in all cases within 48 hours of therapy. There was one case of HELLP syndrome in the 24th week of pregnancy. She developed eclampsia as her third plasma exchange was performed [19].

The application of plasmapheresis/plasma exchange therapy during pregnancy in the following 30 years produced no encouraging results.

More recently, the use of extracorporeal apheresis was reported with success from Thadhani et al. in patients with preeclampsia and elevated levels of sFlt-1 levels [20,21]. The authors described a treatment that reduces circulating sFlt-1 levels. They treated first 8 women with whole blood apheresis and then 11 patients with plasma-specific dextran sulfate column.

There is mounting evidence about the involvement of sFlt-1 in the pathogenesis of human preeclampsia [22]. Other mediators like soluble endoglin show an important role especially in the pathogenesis of HELLP Syndrome [8,22].

Thadhani et al. were not able to refute the possibility that the removal of other factors may also confer benefit to the patients. The use of ligand-specific apheresis columns (eg. configured with anti-sFlt-1 antibody or VEGF) could determine the relative contribution of sFlt-1 depletion versus depletion of other potential mediators of preeclampsia in further studies.
explained this by the renal endothelial cell damage implicated in the pathomechanism of gestational hypertension. The time required to heal this lesion may be the reason for the delayed response to the therapy.

Unfortunately, we failed to prolong pregnancy until the baby could survive after the delivery. An explanation for the limited effect may be the progression of endothelial damage which cannot inhibit by plasmapheresis. However, considering the very early onset of the HELLP syndrome and a prolongation of pregnancy of 20 days in total, plasmapheresis appears to be a promising and safe therapeutic approach to prolong pregnancy in early onset preeclampsia as well as in early onset HELLP syndrome.

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Disclosure of Interests

No author has any conflict of interest to disclose.

Contribution to Authorship

All authors were involved in the conception of the work as well in the critical revision. The first author also wrote the paper.

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