Parametrial hematoma following fetal craniotomy and curettage in intrauterine fetal death: a case report

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ABSTRACT

Background: Parametrial hematoma is a collection of blood located in the parametral area, which is a type of hematoma that can occur in the pelvic cavity. Postpartum hematoma is a rare but life-threatening complication of childbirth. Common risk factors to developing parametrial hematomas include multiple pregnancies, traumatic deliveries, operative vaginal delivery, prolonged labour, manual removal of placenta, inadequate hemostasis at Caesarean section, pre-eclampsia, and anticoagulation therapy. We reported a case of parametrial hematoma post-craniotomy and curettage of a fetus with intrauterine fetal death (IUFD) in a 28-year-old pregnant woman 24 weeks into her fourth pregnancy.

Case report: A pregnant woman with 24 weeks gestation age came to emergency room with complaints of abdominal pain and bloody discharge without clear fluids 9 hours prior. She was diagnosed with preterm delivery and was given tocolytic. The following day, ultrasound examination was done and found fetal heart rate (FHR) was not found, suggesting an intrauterine fetal death (IUFD). Termination was carried out with oxytocin induction but due to maternal exhaustion, pain, and lack of cooperation, a craniotomy was done in operating room followed by curettage. Twenty-four hours after curettage, patient complained of an acute lower right abdominal pain and ultrasound showed a complex mass in right adnexa measuring 8 x 8 cm, suggesting a right adnexal hematoma with a differential diagnosis of a right tubo-ovarian abscess. The patient's haemoglobin was found to decrease to 6.0 g/dl. A laparotomy was performed and a hematoma was found in the right parametrium with active bleeding.

Conclusion: Parametrial hematoma is a rare disease that can occur due to trauma (in labor) or spontaneously due to abnormalities of the uterine arteries that supply blood to the uterus. The patient present in this case report had acute abdominal pain with decreased haemoglobin without signs of bleeding after an operative vaginal birth which may or may not be the cause of the parametrial hematoma due to limitations of examination on the patients. Further observation of similar cases will be required to determine the association between parametrial hematoma and operative vaginal birth.

Keywords: Parametrial hematoma, operative vaginal birth, fetus.

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INTRODUCTION

Concealed postpartum haemorrhage is a case that requires high suspicion and prompt management to reduce morbidity and mortality. Concealed postpartum haemorrhage can occur intrauterine, intra-abdominal, or within a hematoma. Postpartum hematoma is a rare but life-threatening complication of childbirth. Literature reviews have revealed that the incidence of puerperal hematoma varies from 1:309 to 1:1500 deliveries. Incidents can occur immediately after delivery or during the puerperal period. Puerperal hematomas are mainly located on the vulva, vagina, broad ligament and rarely but possible happen in the retroperitoneal area. A pelvic hematoma is a form of postpartum hematoma, which is a collection of blood located in the pelvic peritoneal cavity due to laceration of blood vessels after childbirth or gynaecological procedures. In the pelvic cavity, hematomas can occur in the ureterovesical space, rectouterine pouch (Douglas’ cavity), or in the paravesical, parametrial, paracervical, paravaginal, or pararectal areas. Retroperitoneal puerperal hematoma which occurs secondary to lacerations of the uterine artery or vessels of the broad ligament is the least common hematoma encountered with an estimated incidence of 1/3500 to 1/20,000. The most common locations of hematomas are the vulva, vaginal/paravaginal and retroperitoneal areas resulting from uterine artery injury, and
CASE REPORT

A pregnant woman came to emergency room (ER) with complaints of abdominal pain 9 hours prior. This is the patient’s fourth pregnancy with a gestational age of 24 weeks. The patient complains a bloody discharge without amniotic fluid. On general examination, Glasgow Coma Scale (CGS) was 15 and vital signs were normal. No abnormalities were found on physical examination. On obstetric examination, the uterine fundal height was 18 cm and the estimated fetal weight was 775 grams. On Leopold’s examination, the fetus was palpated with cephalic presentation. Uterine contraction frequency was 2x/10 seconds for 30 seconds, with a fetal heart rate (FHR) of 150x/minute. A vaginal tocher was done and found 2 cm of opening and negative Litmus test. The patient was diagnosed with G P A, with gestational age of 24 weeks, with premature delivery. The patient was treated with tocolytic until uterine contractions disappear.

Three weeks earlier, the patient had visited ER due to abdominal pain, and was given tocolytic therapy, and was discharged after the condition was under control. At that time, the pregnancy condition was good, the gestational age was 20-21 weeks, but the amniotic fluid was found low with an amniotic fluid index (AFI) of 4.7 cm. The cause was not clear. The patient was given oral therapy with aspirin 80mg once a day and was advised to return for a follow-up in the next 2 weeks. The patient has an obstetric history of her first and second children born spontaneously at term with birth weights of 2700 grams and 2400 grams, then a spontaneous abortion in her third pregnancy 3 years ago. The patient used an IUD contraceptive which was removed 8 months ago.

The patient was evaluated the next day to find normal vital signs. On obstetric examination, uterine fundal height was at umbilical level, head presentation, weak uterine contraction, and fetal heart rate (FHR) were not found. On examination, vaginal tocher revealed an opening of 3 cm, membrane was not palpable and dry, and the head was palpable with the transverse sagittal suture with Hodge 1 descent. Ultrasound examination was performed with the results: biparietal diameter (BPD): 6.17 cm ~ 25 mg, head circumference (HC): 22.99 cm ~ 25 mg, abdominal circumference (AC): 20.12 cm ~ 24 mg, estimated fetal weight (EFW): 795 gr, fetal heartbeat (FHB): (–), Spalding sign (–), AFI: 2.1 cm. Blood examinations showed Hemoglobin (Hb) of 7.5 g/dl, white blood cells (WBC) of 13.36 x 10^9/l, and normal coagulation factors. The patient was diagnosed with G P A, 24 weeks of gestation with IUFD, anaemia, and severe oligohydramnios.

Termination of pregnancy was carried out with oxytocin induction. During the observation period, the patient reported feeling pain with visual analogue scale (VAS) around 8-9. The uterine contraction was found inadequate, 2x/10 seconds for 10-15 seconds. A vaginal tocher was done and found an opening of 9 cm, 90% effacement, and palpable left front crown with a Hodge 2 descent. The patient was found lack cooperation with exhaustion and inadequate pushing. A craniotomy was performed on the fetus in the operating room under intravenous anaesthesia followed by curettage to remove any remnants of conception.

After curettage, ultrasound examination was performed with the results: thin uterine endometrial line, no visible intrauterine hyperechoic shadows, normal right and left ovaries, a complex mass is seen in the right adnexa measuring 8 x 8 cm. In conclusion, ultrasound showed a right adnexal hematoma with a differential diagnosis of a right tubo-ovarian abscess. The patient was given antibiotics and analgesics and further observation in the room.

Twenty-four hours post curettage, the patient complained of pain in the lower right abdomen. Vital signs showed blood pressure of 110/80 mmHg, heart rate of 80 bpm, respiratory rate of 18x/minutes, and oxygen saturation of 99%. Abdominal examination revealed tenderness and rebound tenderness in the lower right area. On vaginal examination, no openings were found on vaginal tocher, there was no clear slinger pain in the portio, a mass of 6 x 6 cm was palpable in the right adnexa with pain. In contrast, no abnormalities were found in the left adnexa. Blood examination results showed Hb of 6.0 g/dl. The patient was diagnosed with post-curettage acute abdomen and anaemia, with a differential diagnosis of right adnexal haemorrhage and right tubo-ovarian abscess. A laparotomy was planned, considering there were acute abdominal pain and a decrease in haemoglobin.

Laparotomy was performed with transfusion of 300 cc of PRC. During the procedure, the appendix and uterus looked normal, a hematoma was encountered in the right parametrium measuring 10 x 10 cm. The peritoneum is opened and 300 cc of blood clots were found. No active bleeding was present. The cavity of the former blood clots is filled with sponges and a drain is installed. One day after the laparotomy, the Hb level was 9.5 g/dl post-
transfusion of 600 cc PRC, the fluid in the drain was less than 10 cc. The drain is then removed and the patient is allowed to go home.

DISCUSSION

We report the incidence of parametrial hematoma after a fetal craniotomy and curettage at 24 weeks of pregnancy with IUFD. The diagnosis is based on signs and symptoms, physical examination, supporting laboratory tests, and ultrasound. The hematoma was found and the clot was evacuated during laparotomy surgery. No active bleeding was present.

Hematoma can occur in the parametrium but the incidence is relatively small. The parametrium is a fibrous and fatty connective tissue surrounding the uterus, separating the supravaginal portion of the cervix from the bladder. In the parametrium, there are uterine arteries and ovarian ligaments and are loose connective tissue so that when bleeding occurs in the uterine arteries, they form clots and are accumulated in the parametrium.

Looking at the chronology of the patient where 3 weeks before admission to the hospital the patient experienced contractions and oligohydramnios (AFI 4.7 cm) without a clear cause, it was suspected that there was a uterine artery blood supply disorder that caused a lack of blood supply to the fetus, decreased urine production and even fetal death. At that time, the results of the ultrasound examination did not show any placental abnormalities. Other abnormalities were suspected, but limitations of the patient's health insurance made it difficult for further exploration, such as the use of Doppler ultrasound. It was possible that at that time there was already a uterine artery leak, maybe even a small parametrial hematoma had occurred, but the ultrasound examination was not clear due to the large uterus with the fetus still in it.

In this case, it is suspected that the parametrial hematoma occurred spontaneously and not due to curettage because there was no uterine laceration after the procedure and at the time of laparotomy the uterus was normal and intact, but delivery and curettage may aggravate the hematoma. This is caused by the shrinking (involution) of the uterus that pulls the uterine artery medially so that the bleeding becomes heavier and within 24 hours after the procedure, acute abdominal symptoms occur. This case differs from broad ligament hematoma, which reported that 50% of broad ligament hematoma can be recognized immediately after delivery because these acute hematomas are generally caused by trauma during delivery.

CONCLUSION

Parametrial hematoma can occur due to trauma (in labor) or spontaneously due to abnormalities of the uterine arteries that supply blood to the uterus. Although rare, postpartum hematomas should be considered in puerperal patients presenting with lower abdominal pain following normal vaginal delivery. A hematoma should be considered if there is a significant decrease in hemoglobin (Hb) while there are no signs of bleeding. If oligohydramnios develops without an apparent cause, the possibility of uterine artery bleeding should be considered. Rapid clinical diagnosis and appropriate management are essential to prevent severe hematoma and reduce patient morbidity. Ultrasonography is an important examination procedure to reach a precise diagnosis.

CONFLICT OF INTEREST

All author declares that there is no conflict of interest regarding publication of the case report.

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AUTHOR CONTRIBUTION

All authors had contributed to the manuscript writing and agreed for the final version of the published case report.

ETHICAL ASPECT

The patient had received signed written informed consent regarding publication of their medical data in medical journal as a case report.

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