The effect of uterine artery ligation in patients with central placenta previa: a randomized controlled trial

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Abstract

Background: Placenta previa is major obstetric surgical risk as it is associated with higher percentage of intraoperative and postpartum hemorrhage (PPH), increased requirement of blood transfusion and further surgical procedures. The current study aimed to evaluate uterine artery ligation prior to uterine incision as a procedure to minimize blood loss during cesarean section in patients with central placenta previa.

Methods: One hundred and four patients diagnosed with central placenta previa antenatally and planned to have elective caesarean section were recruited from the antenatal clinic at Minia Maternity University hospital. Patients were randomly allocated into either ligation group or control group.

Results: Both groups were similar regarding demographic features and preoperative risk factors for bleeding. The intraoperative blood loss was significantly lower in the ligation group as compared with the control group (569.3 ± 202.1 mL vs. 805.1 ± 224.5 mL respectively, \( p = 0.002 \)). There was a significant increase in the requirement for blood transfusion in the control group as compared with the ligation group (786 ± 83 mL vs. 755 ± 56 mL respectively, \( p = 0.03 \)). Three cases in the control group required further surgical interventions to control intraoperative bleeding, while no cases in the ligation required further surgical techniques and that was statistically significant (\( p = 0.001 \)).

Conclusion: Uterine artery ligation prior to uterine incision may be a helpful procedure to minimize intraoperative and postpartum blood loss in cases with central placenta previa.

Trial registration: Retrospectively registered in ClinicalTrials.gov Identifier: NCT02002026- December 8, 2013.

Keywords: Central placenta previa, Uterine artery ligation, Cesarean section

Background

Placenta previa is a potentially serious obstetric complication where the placental tissue abnormally lies within the lower uterine segment [1, 2]. The exact pathophysiology of this serious condition is not exactly known. However, uterine scarring is a potential risk factor. Other risk factors for placenta previa include advanced maternal age, high parity, history of placenta previa and congenital uterine malformations [3–6].

The prevalence of placenta previa is estimated to be 5.2 per 1000 pregnancies. However, there is evidence of regional variation [7]. In Minia maternity University hospital where the study was conducted, data from labor ward registry showed that 5% of caesarean deliveries were performed due to placenta previa and its variants. This rate is expected to rise in the coming years due to high rate of caesarean deliveries and subsequently more pregnancies with uterine scarring.

Placenta previa is associated with higher incidence of intraoperative bleeding and postpartum hemorrhage (PPH), need for blood transfusion and further surgical procedures like devascularization and emergency hysterectomy.
Methods
Project no.: MUH201310127
The study protocol was approved by scientific ethical committee of the Department of Obstetrics and Gynecology, Faculty of Medicine, Minia University in September 2013. The study procedure was explained to all eligible patients. All patients signed informed consent that included their agreement to participate in the study.

This study is a randomized controlled study including 104 patients diagnosed with central placenta previa antenatally and were planned to have elective cesarean section. Patients were recruited from attendees of the antenatal clinic at Minia Maternity University Hospital at the period between January 2014 and December 2016.

Patients were randomly allocated into study or control group. The random allocation was based on computer-generated random numbers sealed in consecutively numbered opaque envelopes that were picked up by the patients outside the operating theatre. As major outcomes were not patient-dependent or patient-reported, the patients and the surgeons were not blind to group allocation.

We included patients diagnosed with central placenta previa diagnosed with two-dimensional ultrasound scan at 18 weeks gestation and remained so till the time of planned cesarean delivery. Central placenta previa was defined as placental localization in the lower uterine segment either anteriorly or posteriorly. Laterally situated placenta was not considered “central”. We excluded patients with: a) known bleeding disorder, b) patients with hypertensive disorders or developed preeclampsia (PET) during the study, c) patients who had antepartum hemorrhage (APH) and delivered by emergency cesarean section, and d) patients with anterior placenta previa that were diagnosed antenatally with color Doppler ultrasound and MRI to have placenta accreta. These criteria were designed prior to initiation of the study.

Thorough assessment of the risk factors associated with each case was done. Patients were followed up with regular ultrasound scans every 2 weeks to ensure fetal wellbeing and placental localization till the time of planned delivery. Deliveries were planned to take place between 36 and 38 weeks according to each case situation. Preoperative hemoglobin (Hb) assessment was done and four to six blood units were cross matched for each case before planned cesarean section. Each cesarean section was performed by two consultant obstetricians with experience in operative management of placenta previa and placenta accreta. The two consultants were assisting each other in each case to ensure uniformity of the procedure in all patients. Preoperative hemoglobin level was checked 24 h after the CS (day 1) or after at least 6 h from the blood transfusion. We adhered to the CONSORT guidelines Fig. 1.

Sample size calculation
Sample size was calculated to prevent type II error. The average intraoperative blood loss in cases of central placenta previa at the hospital where the study was conducted was estimated to be 800 ml (figure obtained from the hospital audit report for the year preceding the trial). To be of clinical significance, it was assumed that uterine artery ligation prior to uterine incision should reduce intraoperative blood loss by 50%. Based on these data, we would need to study 27 patients in each arm to be able to reject the null hypothesis that the rates for study and control groups are equal in intraoperative blood loss at a probability of 80%. The type one error probability associated with this test for the null hypothesis is 0.05. To compensate for patients’ withdrawal or cases in which the procedure cannot be performed, we recruited 35 patients in each arm.

Procedure
- Skin incision through Pfannenstiel approach and anterior abdominal wall layers were incised separately.
- The loose peritoneum covering the lower uterine segment is dissected to expose the lower uterine segment and mobilize the urinary bladder downwards.
- Uterine artery ligation was performed by grasping the broad ligament on each side with thumb anterior and the index finger posterior lifting the base below the site uterine incision; the uterine artery was singly ligated with No. 1 vicryl suture. Uterine vessels were ligated and not damaged through inclusion of myometrium. The procedure was then repeated on the other side.
- A curvilinear transverse lower uterine segment incision was performed as usual. Higher incisions
were performed in cases where the traditional incision was expected to be directly through the placenta.
- Delivery of the baby and placenta.
- Closure of the uterine incision in 2 layers with no. 1 vicryl suture.
- Closure of the anterior abdominal wall in layers.

In the control group, lower segment caesarean section was performed in the classic way.

Assessment of intraoperative blood loss
The intraoperative blood loss was measured using the alkaline hematin method [12]. All the blood-stained swabs, diapers and pads and the contents in the drainage bottles were collected, put in a plastic bag and blended with 5% NaOH solution. The plastic bag was then transferred to the Stomacher Lab Blender (Model 3500, Seward Laboratories, London, UK) and processed for few minutes to extract hemoglobin. A portion of the fluid was collected and diluted with 5% NaOH solution. The concentration of alkaline hematin was obtained by assay in a spectrophotometer at 546 nm with the appropriate NaOH as a blank. The intra-assay coefficient for analyzing the concentration is 1%. The blood loss was then calculated using the patient’s preoperative hemoglobin as a reference.

Outcome measures
The primary outcome measures of the study were:
- The amount of intra-operative blood loss.
- The change in pre and post-partum hemoglobin.

The secondary outcome measures were:
- The need for blood transfusion.
- The need for further surgical intervention to control intraoperative bleeding.
- The operative time.
**Statistical methodology**

Statistical analysis was performed using the Statistical Package for Social Science (SPSS Inc., NY) version 21 for Microsoft Windows. Data was described in terms of mean ± SD (standard deviation) for continuous variables and frequencies (number of cases) and percentages for categorical data. Independent Student's t-test was used to compare quantitative variables and Chi square test was used to compare categorical data. A $p$ value < 0.05% was considered significant.

**Results**

We initially recruited 140 patients in this study. In the study group, 10 cases were withdrawn from the study (8 developed APH and had emergency CS and 2 developed PET). In the control group, 14 cases were withdrawn (10 cases developed APH and had emergency CS and 4 cases developed PET).

**Difficulties during the procedure**

In the study group, 12 cases were excluded intra-operatively due to difficulty in performing the procedure as:

1. Extensive varicosities over the lower uterine segment (LUS) in 6 cases
2. Extensive adhesions between the urinary bladder and LUS in cases with repeat cesarean sections in 2 cases
3. Adhesions between the colon and the back of the uterus or the broad ligament in 2 cases
4. Fetal head compressing over the lower uterine segment making fetal head injury possible during the procedure in one case,

In these cases, the procedure was not done, and patients were excluded from the final analysis. However, these cases were analyzed initially within the ligation group. Results remained significant by comparing 56 patients in the control group versus 60 patients in the ligation group (Additional file 1: Table S1, Additional file 2: Table S2, Additional file 3: Table S3 and Additional file 4: Table S4).

At the end, 104 patients reached the final analysis; 48 patients in the study (ligation) group and 56 patients in the control group. Study flow chart is shown in Fig. 1.

There was no statistically significant difference between the study and control groups regarding the demographic features and risk factors associated with placenta previa as shown in Table 1. The ultrasonographic features of the placentae in the study population are summarized in Table 2.

Patients in the ligation group had higher postoperative hemoglobin, which was statistically significant ($p < 0.0001$), shorter operative time that was not significant ($p = 0.2$). The intraoperative blood loss and requirements for blood transfusion were significantly lower in the ligation group compared with the control group ($569.3 ± 202.1$ mL vs. $805.1 ± 224.5$ mL, $p < 0.0001$ and $755 ± 56$ mL vs. $786 ± 83$ mL, $p = 0.03$ respectively). Three cases in the control group required further surgical interventions to control intraoperative bleeding; two cases of internal iliac artery (IIA) ligation and one case of supravaginal hysterectomy. In spite that patients with anterior placenta previa that were diagnosed antenatally with color Doppler and MRI to have placenta accrete were excluded from the study, one patient in the control group was discovered intra-operatively to have placenta accrete and that was the case in whom supravaginal hysterectomy was performed. No cases needed further surgical interventions in the study group. Two cases developed PPH in the control group compared to no cases in the ligation group (3.8% vs. 0%, $p = 0.02$). There were three cases of bladder injuries; one in the ligation group and two in the control group. Bleeding from varicosities over the bladder

| Table 1 Characteristics and risk factors in the study population |
|---------------------------------------------------------------|
| ligation group ($n = 48$) | control group ($n = 56$) | $p$ value |
|---------------------------|---------------------------|-----------|
| Maternal age (years) | $33.5 ± 4.8$ | $34.1 ± 4.7$ | 0.8 |
| BMI (kg/m²) | $28.9 ± 4.7$ | $29.2 ± 5.1$ | 0.7 |
| Occupation: | | | |
| ➢ housewife | 28 | 38 | 0.6 |
| ➢ nonprofessional | 14 | 12 | |
| ➢ professional | 6 | | |
| Residence: | | | |
| ➢ Rural | 34 | 42 | 0.6 |
| ➢ Urban | 14 | 14 | |
| Smoking | | | 0.5 |
| Parity | $4.1 ± 0.9$ | $4.3 ± 1.1$ | 0.7 |
| Previous uterine surgery: | | | |
| ➢ Caesarean section | 20 | 23 | 0.6 |
| ➢ Dilatation and curettage | 8 | 7 | |
| ➢ Myomectomy | 2 | 1 | |
| ➢ Resection of placenta | 0 | 1 | |
| ➢ Resection of intrauterine adhesions | 0 | 2 | |
| ➢ Repair of uterine rupture | 2 | 2 | |
| ➢ B-Lynch suture | 0 | 1 | |
| Previous placenta previa: | | | |
| ➢ Minor | 6 | 7 | 0.5 |
| ➢ Major | 4 | 5 | |
| Manual removal of placenta | 2 | 1 | 0.6 |
| Placenta previa and previous uterine surgery | 8 | 12 | 0.7 |
| Pregnancy with assisted conception | 2 | 4 | 0.5 |

Data are presented as mean ± SD or frequency and percentages.
Regional anesthesia was used in all cases at the beginning. In the control group, six cases were converted to general anesthesia (GA) as spinal anesthesia worked off and additional surgical procedures were required and consumed longer time (two cases of internal iliac artery “IIA” ligation and one case of supravaginal hysterectomy). In these three cases, placenta was anterior. No cases in the study group required conversion to GA (0% vs. 23.1%, \( p = 0.001 \)).

There was no significant difference in the experience of anesthetist between the two groups. Anesthetic details are summarized in Table 4.

The mean gestational age at time of CS was \( 36.2 ± 1.1 \) weeks in the study group and \( 36.3 ± 0.9 \) weeks in the control group (\( p = 0.7 \)). There was no significant difference between the two groups regarding the perinatal outcome difference as shown in Table 4.

Discussion
To our knowledge, it is the first study to assess the technique of uterine artery ligation in patients with central placenta previa. We recruited patients diagnosed with central placenta previa at 28 weeks’ gestation by 2D ultrasound. We used the alkaline hematin test to allow objective assessment of the intraoperative blood loss.

The new procedure evaluated in this study was found to minimize the intraoperative blood loss and shorten the operative time. Patients in the study group had

Table 3 Outcome measures in the ligation and control groups

|                         | Ligation group (\( n = 48 \)) | Control group (\( n = 56 \)) | \( p \) value |
|-------------------------|-------------------------------|-------------------------------|--------------|
| Preoperative Hb (g/dL)  | 11.1 ± 0.61                   | 11.2 ± 0.6                   | 0.6          |
| Postoperative Hb (g/dL) | 10.2 ± 0.34                   | 9.3 ± 0.56                   | 0.0001*      |
| Intraoperative blood loss (mL) | 569.3 ± 202.1 | 805.1 ± 224.5 | 0.0001*      |
| Operative time (min)    | 54.2 ± 11.2                   | 57 ± 10.9                    | 0.2          |
| Amount of blood transfusion (mL) | 755 ± 56          | 785 ± 83                     | 0.03*        |
| Further surgical interventions: |                        |                              |              |
| ➢ IIA ligation          | 0 (0%)                        | 2 (3.6%)                     | 0.001*       |
| ➢ Supravaginal hysterectomy |                        | 0 (0%)                        |              |
| Urinary bladder injury  | 1 (2.1%)                      | 1 (1.8%)                     | 0.2          |
| Bleeding from varicosities over bladder surface | 2 (4.2%) | 3 (5.4%) | 0.6 |
| Postpartum hemorrhage (n) | 0 (0%)                        | 2 (3.6%)                     | 0.02*        |

12 cases were excluded from ligation group in the final analysis due failure of the procedure. Results remained significant by comparing 56 patients in the control group versus 60 patients in the ligation group.

Data is presented as mean ± SD or frequency and percentages

Table 4 Anesthetic details of CS and Perinatal outcome in the study population

|                         | Ligation group (\( n = 48 \)) | Control group (\( n = 56 \)) | \( p \) value |
|-------------------------|-------------------------------|-------------------------------|--------------|
| Type of anesthesia:     |                               |                               |              |
| ➢ Spinal                | 38 (79.1%)                   | 43 (76.8%)                   |              |
| ➢ Epidural              | 10 (20.9%)                   | 13 (23.2%)                   |              |
| ➢ GA††                  | 0 (0%)                        | 3 (5.3%)                     |              |
| Gestational age at time of CS (weeks) | 36.2 ± 1.1               | 36.3 ± 0.9                   | 0.7          |
| Birth weight (gram)     | 2875.6 ± 253.5               | 2976.5 ± 265.2               | 0.5          |
| Apgar score at 5 min:   |                               |                               |              |
| ➢ 1–2                   | 2 (4.2%)                      | 2 (3.6%)                     | 0.7          |
| ➢ 3–6                   | 12 (25%)                      | 16 (28.6%)                   | 0.6          |
| ➢ ≥7                    | 34 (70.8%)                    | 38 (67.8%)                   | 0.8          |
| Neonatal outcome:       |                               |                               |              |
| ➢ Early neonatal death  | 1 (2.1%)                      | 1 (1.8%)                     | 0.88         |
| ➢ NCU admission         | 4 (8.4%)                      | 4 (7.2%)                     | 0.9          |
| ➢ Hospital discharge    | 44 (91.6%)                    | 51 (91%)                     | 0.91         |

Data is presented as frequency and percentages or mean ± SD

GA: general anesthesia
††: Conversion from spinal to general anesthesia as spinal worked off (2 cases of IIA ligation and 1 case of supravaginal hysterectomy)

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higher postoperative hemoglobin and lower requirement for blood transfusion. In addition, none of the patients underwent the procedure needed further surgical interventions to control intraoperative bleeding, while three patients in the control group needed another surgical intervention; two IIA ligation and one case needed supravaginal hysterectomy.

The procedure was not easy in all cases and was not possible in some patients as those with extensive adhesions between the LUS and the urinary bladder, extensive varicose veins in the LUS. In these cases, the procedure was not done as we believed there would be a great risk of urinary bladder injury or severe bleeding from varicose veins. In some occasions, the fetal head was compressing LUS, and the ligation was not done.

Ligation of Internal iliac artery, used to be performed to overcome massive pelvic hemorrhage, is not the technique of choice for control of atonic PPH due to placenta previa. Apart from its efficacy it requires more time for dissection and effort for training [13]. In a comprehensive research for 30 years, O’Leary had suggested the procedure of uterine artery ligation as an efficient and alternate procedure to internal iliac artery ligation although in his trial, 10 out of 265 cases failed to respond [14]. In another excellent study concerning massive PPH due to uterine atony. The authors concluded that arterial embolization as effective as uterine artery ligation with success rate approaching 100%. Embolization can be performed after vaginal delivery in stable patients but it needs expertise and facilities [15].

The strengths of this study are the nature of the study (RCT) and reporting a novel technique in management of a serious obstetric condition as central placenta previa with potential for implication in clinical practice. The limitation of the study is the relatively small number of patients included.

Conclusion
Uterine artery ligation prior to uterine incision could be an effective method to reduce the intraoperative blood loss in patients with central placenta previa undergoing elective CS. Larger studies are required to reach a firm conclusion about the procedure.

Acknowledgements
The authors are grateful to the entire medical, nursing and laboratory staff at Minia Maternity University Hospital for their help and cooperation throughout the research work.

Availability of data and materials
The datasets used and analyzed during the current study are available from the corresponding author on reasonable request. All data generated & analyzed during this study are included in this published article.

Authors’ contributions
ASS: Design the study methodology, diagnosis of cases, operative management, data collection, results preparation, and manuscript writing. AEM: Design the study methodology, diagnosis of cases, operative management, data collection, results preparation, and manuscript writing. HMK: Design the study methodology, operative management, diagnosis of clinical cases program, and data collection. HAB: Design the study methodology, operative management, diagnosis of cases. HM: Design the study methodology, diagnosis of clinical cases program, and data collection. RRE: Design the study methodology, diagnosis of clinical cases program, and data collection. AGA: Design the study methodology, operative management, data collection, results preparation and manuscript writing. MAZ: Design the study methodology, diagnosis of cases, operative management, data collection, results preparation and manuscript writing. HHK: Design the study methodology, diagnosis of clinical cases program, and data collection. We insist us in ethical consideration, help in reviewing the work, was involved in revising the manuscript and provide final approval of the version to be published. All authors read and approved the final manuscript.

Ethics approval and consent to participate
The study protocol was approved by scientific ethical committee of the Department of Obstetrics and Gynecology, Faculty of Medicine, Minia University in September 2013. Approval was ascertained from the Institutional Review Board of the Faculty of Medicine, Minia University in October 2013. All procedures performed were in accordance with the ethical standards of the 1964 Helsinki declaration and its later amendments or comparable ethical standards. All patients signed informed consent that include their agreement to participate in the study.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Received: 30 November 2017 Accepted: 21 August 2018 Published online: 29 August 2018

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