**Introduction**

We report here on an unusual case of placenta percreta. Placentation disorders (placenta accreta, increta, and percreta) occur in around 2% of all births [1]. The incidence of placenta percreta reported in the literature ranges between 1 : 7000 [2] and 1 : 2500 [3] pregnancies. Due to increasing numbers of C-sections, however, over the last few years the incidence appears to be on the rise.

After a diagnosis of placenta percreta has been made ante partum, there are four main approaches for clinical management:

1. C-section with hysterectomy
2. Planned in situ retention of the placenta percreta with embolisation of the uterine artery
3. Planned in situ retention of the placenta percreta with subsequent methotrexate therapy
4. Planned in situ retention of the placenta percreta with expectant management and close monitoring

The main criterion for conservative management of placenta percreta is the wish of the patient to have more children [1]. Moreover, in cases of placenta percreta with invasion of the bladder, conservative management results in fewer operative complications, such as fistula formation or injury to the ureter, both of which are associated with high maternal morbidity [3]. Complications of conservative management of placenta percreta described in the literature include bleeding, infection (endometritis, wound infection, peritonitis, pyelonephritis, uterine necrosis), sepsis and septic shock, fistula formation, thrombosis, pulmonary embolism, pulmonary oedema, and the side-effects of methotrexate therapy [3]. Acute renal failure has only been described in one case with methotrexate injection into the umbilical cord and was considered an acute side-effect of methotrexate therapy [3]. Acute renal failure has only been described in one case with methotrexate injection into the umbilical cord and was considered an acute side-effect of methotrexate therapy [3].

**Abstract**

A 22-year-old gravida II, para I, with a twin pregnancy was diagnosed with placenta praevia totalis et percreta in 25 GW. After consideration of various modes of delivery a C-section was performed with retention of the placenta percreta in situ when vaginal bleeding occurred in 28 GW. Following 8 dosages of methotrexate given on an outpatient basis, the patient suddenly developed acute renal failure necessitating dialysis. This was due to a bilateral renal cortical necrosis after disseminated intravasal coagulation based on a massive accumulation of trophoblastic tissue.

**Zusammenfassung**


**Bibliography**

DOI http://dx.doi.org/10.1055/s-0032-1327941

Geburtsh Frauenheilk 2012; 72: 1033–1035 © Georg Thieme Verlag KG Stuttgart · New York · ISSN 0016-5751

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**Key words**

- twin pregnancy
- placenta praeavia
- placenta percreta
- renal failure

**Schlüsselwörter**

- Mehrlingsschwangerschaft
- Placenta praeavia
- Placenta percreta
- Niereninsuffizienz
67% of patients managed conservatively became pregnant again [1].

Case Report

A 22-year-old gravida II, para I, with a previous history of secondary C-section and repeat pregnancy in rapid succession, presented with a dichorial diamniotic twin pregnancy and was diagnosed with placenta praevia totalis in the 21st gestational week (GW). The patient was admitted to our hospital in the 24 + 1 GW with acute vaginal bleeding. In addition to placenta praevia totalis, sonographic imaging was suspicious for placenta percreta of the 1st twin with the placenta of the 2nd twin immediately adjacent on the uterine wall (Fig. 1). This suspicion was confirmed by cystoscopy. During the patient’s stay in hospital, various modes of delivery including C-section followed by hysterectomy or retention of the placenta percreta in situ followed by methotrexate therapy were discussed with the patient and her partner. Based on the overall combination of findings they opted to leave the placenta percreta in situ with subsequent methotrexate therapy.

After the recurrence of vaginal bleeding and the onset of labour, secondary C-section was performed in the 27 + 2 GW and the twin placenta were left in situ. As the placenta of the 2nd twin was directly adjacent to the placenta percreta, it was considered too risky to remove the 2nd placenta. Two preterm infants were delivered, in differing stages of growth. Both adapted well and continued to develop according to their stage of maturation. The patient received 2 units of packed red blood cells perioperatively. On the 1st postoperative day, methotrexate therapy with the dosage tailored to the patient’s weight (76 mg = 50 mg/m² median BSA) was initiated. Different therapy regimes for methotrexate dosages and application to treat placenta percreta have been described in the literature [2,4,5]. The above dosage chosen by us and the application interval had previously been used successfully in another patient with placenta percreta treated in our hospital and corresponded to dosages given in the British therapy guidelines for the conservative management of tubal pregnancy [6]. On the 13th postoperative day, the patient was discharged home in good general health with further monitoring done on an outpatient basis.

The patient continued to receive a weekly application of methotrexate at the dosages described above; each after routine laboratory investigations such as complete blood count, CRP, and creatinine levels. The patient received a total of 8 × 76 mg doses of methotrexate administered intramuscularly. Throughout this period, levels of β-HCG were controlled regularly and the patient was monitored with regular sonograms. Initially, vascularity of the placenta praevia and percreta regressed well, but as levels of β-HCG in serum stagnated, vascularisation began to increase again (Table 1).

The patient was readmitted to hospital 4 days after the last administration of methotrexate with moderate vaginal bleeding and impaired coagulation (Quick’s value: 56%, partial thromboplastin time [PTT]: 36 s). After a brief time in the intensive care unit where the bleeding was stopped and coagulation parameters improved, the patient was initially transferred to the general ward again, and conservative management of the placenta percreta still appeared possible. Three days later, however, there was strong clinical deterioration with renal failure necessitating dialysis (creatinine increase from 0.5 mg/dl on admission to 3.4 mg/dl after a few days), and incipient pulmonary oedema. The patient also presented with intermittent accompanying symptoms such as diarrhoea, vomiting, icterus, and pancytopenia. After the patient was stabilised in the intensive care unit, secondary hysterectomy was performed with reconstruction of the bladder. Intraoperatively, the patient required mass transfusion and the administration of coagulation factors. Two days after the operation, the patient was transferred to the nephrological intensive care unit for further treatment. Due to the pulmonary oedema she required ventilation for 10 more days. She continued to be anuric and required dialysis.

Subsequently, comprehensive differential diagnostics were done. Based on the accompanying gastrointestinal symptoms described above and the pancytopenia, there was an initial suspicion that the acute renal failure was a consequence of methotrexate intoxication. But the serum methotrexate levels were already below the detection limit a few days after administration of the last dose; moreover, the overall low dosages administered (one

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<th>Date</th>
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tenth of the dosage administered in tumour therapies) and subsequent unremarkable bone marrow aspiration militated against an overdose of methotrexate. Hypovolaemic shock as the cause of acute renal failure appeared unlikely, as vaginal bleeding was quickly stopped after admission to hospital, and the patient’s vital parameters remained unremarkable until she required intubation. Sepsis was excluded based on the constantly stable CRP levels and negative procalcitonin. Haemolytic-uraemic syndrome (HUS) was excluded because of the rapid increase in thrombocytes after the administration of thrombocyte concentrates and the too low numbers of schistocytes; similarly, EHEC and hantavirus serology found no pathogens. Massive preeclampsia was also discussed as a possible cause but was excluded from the differential diagnosis due to the fact that the patient’s blood pressure remained normal, she had no proteinuria and the sFlt/PlGF ratio was unremarkable.

Further diagnostic imaging finally showed bilateral necrosis of the renal cortex on MRT as the cause of the existing renal failure. Based on laboratory findings, which showed a fibrinogen level of < 50 at the time of the dramatic clinical deterioration, the assumption was that pronounced disseminated intravasal coagulation was the cause of the bilateral renal cortical damage. This was most probably triggered by a massive accumulation of trophoblastic tissue from the twin placenta remaining in situ at the time of the recurrence of vaginal bleeding. It is known that extensive necrosis of the renal cortex can occur as a result of a thrombosis of the interlobular artery and the afferent vessels, particularly after an endotoxic shock (in our case through the accumulation of trophoblasts) with disseminated intravasal coagulation [7]. As this pathomechanism appeared to be the most plausible one, the nephrologists in attendance decided against performing invasive renal biopsy.

One year after renal failure the patient permanently requires dialysis and is on the list for kidney transplantation.

**Discussion**

A case like the one presented here of placenta percreta with bilateral renal cortical damage and subsequent chronic renal failure has not been previously described in the literature. We therefore suggest that careful evaluation and detailed diagnostics are important if placenta percreta is left in situ. The infiltration of adjacent organs, the wish of the mother to have more children, and the special course in cases with twin placentas with large amounts of trophoblastic material are all important points to watch. If conservative management is chosen post partum, frequent and regular sonographic monitoring together with monitoring of coagulation parameters and renal function is required.

**Conflict of Interest**

None.

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