Retained placenta after vaginal delivery: risk factors and management

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Abstract: Retained placenta after vaginal delivery is diagnosed when a placenta does not spontaneously deliver within a designated amount of time, variably defined as a period of 18–60 mins. It may also be diagnosed if a patient experiences significant hemorrhage prior to delivery of the placenta. Normal placenta delivery requires adequate uterine contractions, with shearing of the placenta and decidua from the uterine wall and expulsion of the tissue. Thus, retained placenta can occur in the setting of significant uterine atony, abnormally adherent placenta, as with placenta accreta spectrum (PAS), or closure of the cervix prior to placental expulsion. Risk factors for retained placenta parallel those for uterine atony and PAS and include prolonged oxytocin use, high parity, preterm delivery, history of uterine surgery, and IVF conceptions. History of a prior retained placenta and congenital uterine anomalies also appear to be risk factors. Management entails manual removal of the placenta with adequate analgesia, as medical intervention alone has not been proven effective. Complications can include major hemorrhage, endometritis, or retained portions of placental tissue, the latter of which can lead to delayed hemorrhage or infection. Prophylactic antibiotics can be considered with manual placenta removal, though evidence regarding effectiveness is inconsistent. If hemorrhage is encountered, deployment of a massive transfusion protocol, uterine evacuation with suction, and use of intrauterine tamponade, as with an intrauterine balloon, should be initiated immediately. When a separation plane between the placenta and uterus is particularly difficult to create, PAS should be considered, and preparations should be made for hemorrhage and hysterectomy. Patients with risk factors for retained placenta should have a laboratory sample sent for blood type and antibody screening on admission to labor and delivery, and plans should be made for appropriate analgesia and preparations for hemorrhage if a retained placenta is encountered.

Keywords: retained placenta, manual removal of the placenta, postpartum hemorrhage, placenta accreta spectrum

Introduction

Retained placenta after vaginal delivery, which occurs in around 1–3% of deliveries, is a relatively common cause of obstetrical morbidity. This is typically diagnosed when the placenta fails to spontaneously separate during the third stage of labor when a patient experiences excessive bleeding in absence of placenta separation or if there is confirmation of placenta tissue remaining after the majority of the placenta delivers spontaneously.1–3 Placetas that fail to spontaneously separate can be a cause of significant surgical and hemorrhagic morbidity.4,5 Untreated, retained placenta is considered the second leading cause of postpartum hemorrhage (PPH).5,6
Although retained placenta is an obstetrical complication encountered relatively infrequently on the labor and delivery floor, recognizing patient risk factors and understanding management are important steps in mitigating this morbidity.

**Pathophysiology**

Normal placentation begins with blastocyst implantation into the maternal endometrium. In preparation for this implantation, the endometrium develops the decidua under the influence of progesterone and estrogen in early pregnancy. As the blastocyst invades this decidua, the layer of cells forming the surface of the blastocyst develops into the chorionic membrane. Cytotrophoblast cells proliferate from the chorionic membrane and form multinucleated aggregates called syncytiotrophoblast cells. These cells form the placental villi, allowing fetal–maternal interchange between the villi–decidual interaction. With delivery of the infant, both a hormonal cascade and uterine contractions allow for separation of these layers and expulsion of the placenta.7

Retained placenta is generally attributed to one of three pathophysiology. First, an atonic uterus with poor contraction may prevent normal separation and contractile expulsion of the placenta.2,8,9 Second, an abnormally adherent or invasive placenta, as seen with placenta accreta spectrum (PAS), may be incapable of normal separation. Finally, a separated placenta may be trapped or incarcerated due to closure of the cervix prior to delivery of the placenta.2,8–10 Placental hypoperfusion disorders, such as with preeclampsia, and infection have also been proposed as mechanisms for retained placenta, although little is known about the specific mechanism.9,11

**Epidemiology**

Estimates of retained placenta put the incidence at between 0.1% and 3%.5,8 Prospective investigations of retained placenta confirm these estimates, with one study of >45,000 patients showing that overall for all gestational ages, retained placenta happened in about 3% of deliveries, with gestational ages of <26 weeks and <37 weeks having a significantly increased risk of retained placenta requiring manual removal.1 Generally, incidence seems to be higher in developed countries where practices tend toward earlier manual removal of the placenta in the third stage of labor.8,12

**Risk factors**

Many studies have attempted to define risk factors for retained placenta, which are listed in Table 1. Established risk factors include prior retained placenta, preterm delivery, prior uterine surgery, previous pregnancy termination, miscarriage or curettage, grand multiparity (greater than five prior deliveries), and congenital uterine anomalies (often unrecognized prior to delivery).3,5,11

Some studies have suggested that prolonged oxytocin use could be a potentially modifiable risk factor for retained placenta, with one study reporting that oxytocin use for over 195 mins increased the odds ratio of the retained placenta by 2.0, and oxytocin use over 415 mins increased the odds ratio by 6.5.5 It is less clear whether oxytocin is directly involved in placental retention, or if the association is mediated by uterine atony or infection due to prolonged labor.

Placental under perfusion disorders have been implicated as risk factors for retained placenta.11 In a case–control study of all singleton primiparous vaginal deliveries in Sweden between 1997 and 2009, the authors found an increased association between placental under perfusion disorders (such as preeclampsia, small for gestational age, and stillbirth) and retained placenta; however, they could not designate a common pathophysiology.

Some research suggests that women may be predisposed to retained placenta. Retained placenta in a prior delivery appears to be an important risk factor for recurrence. In one study of over 280 women in Denmark, prevalence of retained placenta was found to be consistent with previously reported numbers (approximately 3%) using strict diagnostic criteria. The authors found that in subsequent vaginal deliveries, the risk of recurrence was substantially increased to about 25%.3 There has even been some suggestion that tendency toward retained placenta may even be inherited. In one study, authors used the Swedish Medical Birth Register to identify women with

<table>
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<th>Table 1 Risk factors for retained placenta</th>
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<tr>
<td><strong>Risk factors related to poor uterine contraction</strong></td>
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<tr>
<td>High parity</td>
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<td>Prolonged use of oxytocin</td>
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<td><strong>Risk factors related to abnormal placentation</strong></td>
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<td>History of uterine surgery</td>
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<td>IVF conception</td>
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<td><strong>Other risk factors</strong></td>
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<td>Preterm delivery</td>
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<td>Congenital uterine anomaly</td>
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<td>Prior history of retained placenta</td>
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retained placenta after 1992 whose mothers’ own birth records were also in the Register (after 1973). The authors found that the risk of retained placenta increased if retained placenta had occurred at the mother’s own birth (aOR 1.66, 95% CI 1.52–1.82), at the birth of one of her siblings (aOR 1.58, 95% CI 1.43–1.76), or both (aOR 2.75, 95% CI 2.18–3.46).13

Because of its relationship to PAS, assisted reproductive technologies (IVF or ICSI) have been proposed and studied as an additional risk factor for retained placenta.14 Elenis et al, in a 2015 study from Sweden, looked specifically at oocyte donation IVF and the risk of poor obstetrical outcomes in otherwise healthy women.15 The authors found a positive association between retained placenta and oocyte donation, as well as between PPH and oocyte donation.15 In another 2016 study by Aziz et al, seeking to determine whether or not length of third stage was related to IVF, the authors concluded that cryopreserved embryo transfer (donated or autologous) without controlled ovarian hyperstimulation was not related to longer third stage, but did significantly increase the risk for manual removal of the placenta.16

Morbidity
Retained placenta requiring invasive procedures is associated with obstetrical morbidities. Of arguably greatest significance is the risk of postpartum hemorrhage, with retained placenta the second leading cause of significant and even fatal hemorrhage in the obstetric population.5,17 One group found that the odds ratio related to estimated blood loss exceeding 500 mL, 1000 mL, and 2000 mL with retained placenta, respectively, is as high as 33.07 (95% CI 20.57–53.16), 43.44 (95% CI 26.57–71.02), and 111.24 (95% CI 27.26–454.00).5 In another case–control study of 114 women with manual removal for retained placenta, the authors found that the case group required significantly more blood transfusions (13% in the case group versus 0% in the controls).18 Large cohort studies have confirmed this elevated risk.17

Further research additionally suggests that the longer the third stage of labor, the greater the risk of postpartum hemorrhage.19 A study by Dombrowski et al in 1995 tried to determine gestational age–specific data for the length of the third stage, retained placenta, hemorrhage, and manual removal. The authors found that both manual removal of the placenta and PPH decreased with increasing gestational age, and that the two were related. However, causal association could not be determined.1

If the placenta or pieces of the placenta remain in situ following attempt at manual removal, a patient may require surgical management. In a study of >20,000 patients in Norway, 3% of women requiring manual removal of retained placenta needed additional surgical management with dilation and curettage.17 Another case–control study of 114 women found that cases required more dilation and curettage than controls, although with their study number they could not confirm significance.18 Occasionally portions of the placenta or membranes may remain in the uterus after manual extraction, leading to delayed complications from retained products of conception. These can include delayed postpartum hemorrhage or endomyometritis.

Evidence of infection risk, particularly endometritis, following manual or surgical removal of retained placenta has been inconsistently demonstrated.19 A large 1995 retrospective cohort study at University of Iowa compared over 1000 patients requiring manual extraction after vaginal delivery with those who did not.20 After controlling for confounders, the authors found that manual removal of retained placenta was significantly associated with postpartum endometritis.20 Alternatively, in the large cohort study of >20,000 patients from Norway mentioned above, patients requiring intervention for retained placenta did not show a significantly increased risk of infection, despite varying practices regarding antibiotic administration and timing.17 Other studies have similarly found a relationship but could not prove a significant association between manual removal or surgical placental removal and endometritis.18,21 The discrepancies may in part be due to the lack of rigorous distinction between postpartum fever and true uterine infection.

Diagnosis
Retained placenta is clinically diagnosed when the placenta fails to spontaneously separate during the third stage of labor, with or without active management, or in the setting of severe bleeding in the absence of placental delivery.16,22 The first diagnostic criterion requires a time cutoff, though there is no uniform consensus as to timing for diagnosis of retained placenta in the third stage in the absence of postpartum hemorrhage. Selection of a clinical time definition can be based either on a population curve of observed spontaneous placental delivery times or on a time at which morbidity significantly increases. Thirty minutes have been used as a loose guideline, which comes from a 1991 study by Combs et al.2 The researchers...
found that the third stage had a log-normal distribution, with a mean length of 6.8 minutes, with only 3.3% of deliveries having greater than 30 minutes third stage. This timing has been supported by other studies as well. Interestingly, the authors calculated that the incidence of PPH, transfusion, and dilation and curettage remained constant during this period, increasing only after 30 minutes and plateauing at 75 minutes for both manually and spontaneously delivered placentas. Because PPH incidence did not increase until after 30 minutes, Combs et al recommended this timing for initiation of manual removal of the placenta.

However, this guidance is not uniformly supported. In a subsequent study by Deneux-Tharaux, surveys from 14 European countries exhibited wide variations in wait time prior to manual placental removal, largely by country but also by the hospital. In countries such as Finland and Denmark, obstetricians tended to wait 60 minutes or more prior to manual removal of the placenta, versus in countries such as Spain and France, where providers removed the placenta after 30 minutes. Practices also varied considerably depending on whether or not the patient in question had prior epidural anesthesia. National and worldwide guidelines similarly have no consensus on when to intervene on an undelivered placenta. For instance, the National Institute for Health and Clinical Excellence suggests a wait time of 30 minutes in the United Kingdom prior to manual removal of the placenta, while the World Health Organization guidelines propose a wait time of 60 minutes.

The most significant risk of waiting a prolonged amount of time before removing the placenta is postpartum hemorrhage. In 2005, Magann and colleagues undertook a prospective observational study in which all women delivering vaginally were assessed for PPH. Using receiver operating characteristic curves, the authors showed that 95% of normal placental delivery occurs within 18 minutes, and that a third stage of labor longer than 18 minutes was associated with a significant risk of PPH. The authors followed up this paper in 2012 with a randomized controlled trial assigning vaginal deliveries to manual removal at either 10 or 15 minutes (as opposed to the traditional 30) if the placenta had not yet spontaneously delivered. The findings supported the authors’ initial study, showing that removal at 15 minutes had a significantly greater likelihood of hemorrhage compared to 10 minutes, opening up the discussion on earlier intervention.

At times the bulk of the placenta will deliver spontaneously or manually, but small portions or an accessory lobe may be retained. This may be suspected when the placenta appears fragmented after delivery or when there is ongoing heavy uterine bleeding. In this situation, the uterine cavity may be evaluated with manual exploration or with ultrasound. The utility of ultrasound in this situation has yet to be established, with a focal endometrial mass, particularly with Doppler flow, being the findings of interest. In one study of routine ultrasound immediately after vaginal delivery, the sensitivity for diagnosing retained placental fragments was only 44% with a positive predictive value (PPV) of 58%. An alternate study showed a 75–80% sensitivity of postpartum ultrasound, though the mean time for evaluation was 21 days postpartum, when less blood and decidua are expected to be seen. While immediate ultrasound’s PPV will be higher when there is clinical suspicion of retained POCs, a negative ultrasound should not deter manual or suction curettage when there is a strong clinical suspicion, especially in the setting of hemorrhage.

Management

After delivery of the infant and prior to diagnosis of retained placenta, active management is recommended to facilitate spontaneous placental separation, including oxytocin, controlled cord traction, and uterine massage. These maneuvers have been shown to decrease the risk of postpartum hemorrhage, though it has not been shown that active management will prevent retained placenta.

Once diagnosed, the placenta is usually manually extracted from the uterus. Table 2 lists items that should be readily available if needed during the extraction process. Because this procedure is painful, adequate analgesia should be achieved via epidural, conscious sedation, or general anesthesia prior to an attempt at extraction. Once the patient is comfortable, she should be appropriately positioned in lithotomy. A conical drape, preferably one that is graduated and marked to allow for quantitative blood loss, should be placed under the patient’s buttocks. The operator should make every attempt to wear gown and gloves and maintain sterility, both for personal and for patient protection. The patient’s bladder should be drained. The provider should then use one hand to follow the umbilical cord through the vagina and cervix until the placenta is palpated. If the placenta is separated but not expelled, such as in the case of uterine atony, the tissue can be firmly grasped and brought through the cervix.
Items that should be available for manual placental extraction

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<th>Medications</th>
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<td>Analgesics or sedatives with appropriate monitoring equipment</td>
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<td>Uterotonic agents</td>
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<td>Nitroglycerin</td>
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<th>Patient preparation</th>
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<td>Conical under-buttocks drape for blood collection</td>
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<td>Bladder catheter</td>
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<th>Surgical instruments</th>
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<tr>
<td>Sponge forceps for grasping tissue</td>
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<td>Long curette</td>
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<th>Hemorrhage management</th>
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<td>Intrauterine balloon or uterine packing material</td>
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<td>Availability of crossmatched blood</td>
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<tr>
<td>Massive transfusion protocol</td>
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<tr>
<td>Equipment for quantitative blood loss measurement (volumetric or gravimetric)</td>
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<tr>
<td>Available operating room with anesthesia equipment</td>
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<tr>
<td>Suction curettage equipment</td>
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<td>Laparotomy equipment</td>
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Uterotonic medications, such as oxytocin, methylergonovine, carboprost, or other prostaglandins, should be given to facilitate contraction once the placenta is removed. If placental removal is refractory or only partially successful (ie the placenta or parts of the placenta remain in the uterus), or if bleeding persists despite placental delivery, often the next step is surgical management with curettage. This may be best achieved in an operating room, with optimal access to surgical equipment, analgesia, and patient resuscitation aids, if needed. Suction curettage is generally used, though a sharp curette may be needed to facilitate a separation plane. Access to uterine tamponade supplies with either a large intrauterine balloon or surgical packs should be immediately accessed in the event of hemorrhage. Crossmatched blood products should be made imminently available if placental separation is difficult or blood loss exceeds 1 L, and the care team should attend to uterotonics and attention to coagulopathy as the extraction is performed. Due to the risk of endometritis, routine antibiotics are generally administered just before or shortly after manual removal of the placenta. Prophylaxis can parallel cesarean prophylaxis with a first-generation cephalosporin. Patients who are febrile at the time of extraction should be fully treated for chorioamnionitis with broad-spectrum antibiotics. Despite these guidelines, few studies have been undertaken examining the effectiveness of antibiotics in reducing infectious morbidity. A 2015 systematic review by Chibueze and colleagues attempted to summarize the literature on the efficacy of antibiotics for preventing adverse maternal outcomes related to manual placenta removal following vaginal birth. The authors reported on three retrospective cohort studies examining endometritis and puerperal fever after manual extraction for retained placenta. None of the three studies found evidence to suggest beneficial effects for routine antibiotic use in women undergoing intervention for retained placenta. The authors concluded that further research is required to adequately answer this question. Due to mixed data regarding prophylaxis, as well as the increasing risk of postpartum hemorrhage with prolonged third stage of labor, administration of antibiotics should not delay manual removal of retained placenta.

Occasionally, a portion of placental tissue may remain in the uterus, either knowingly or unbeknownst to the providers. This can present as abnormal bleeding days to weeks after delivery and should be suspected in the setting of a delayed postpartum hemorrhage. Recently, studies have examined the usefulness of hysteroscopic morcellation devices in aiding with retained placenta left in situ postpartum (Figure 1A [before] and B [after]). In a series...
of case reports, Lee and colleagues reported a higher risk of complications with blind curettage compared to hysteroscopic morcellation.\textsuperscript{33} They additionally reported complete resection in 90\% of hysteroscopic cases and reduction of both perforation and intrauterine adhesion risk.\textsuperscript{33} In another randomized control trial by Hamerlynck et al, the authors randomized patients to undergo hysteroscopic resection of retained placenta with either hysteroscopic morcellation versus loop resection with rigid bipolar resectoscopes.\textsuperscript{34} These authors in comparison found that when comparing the two modalities, complete resection was comparatively high in both groups, and intrauterine adhesions were comparatively low.\textsuperscript{34} The one significant difference between the two groups was that the hysteroscopic group had significantly faster operative times.\textsuperscript{34} The ability to perform hysteroscopic removal depends on the amount of active bleeding, with suction curettage often needed when bleeding is heavy.

Other studies have examined alternative, nonsurgical, management for retained placenta, none of which have been successful. In 2012, 99 women in a large teaching hospital in the Netherlands with retained placenta (>60 mins after delivery) were given either 800 mcg misoprostol or placebo orally.\textsuperscript{35} The author’s primary outcomes were number of manual removals of retained placenta and blood loss. The authors found that oral misoprostol reduced neither the need for manual removal nor the overall amount of blood loss. Both groups were observed for additional 45 mins after administration of misoprostol or placebo. While the authors found that 50\% of remaining placentas at 60 mins delivered in the intervening 45 mins, it came at the expense of additional significant blood loss.\textsuperscript{35}

For a time, umbilical vein oxytocin was thought to be a promising alternative or adjunct to manual extraction of the placenta. A 2011 Cochrane Review summarized available data on the subject to assess the use of umbilical vein oxytocin either alone or in conjunction with intravenous oxytocin to reduce the need for manual removal of retained placenta.\textsuperscript{36} While inexpensive and easy to do, the authors found that all well-designed randomized control trials showed no significant effect of umbilical vein oxytocin on retained placenta.\textsuperscript{36}

\section*{Morbidly adherent placenta}

In the unusual event that manual extraction does not result in delivery of the entire or partial placenta, MAP must be considered as an etiology. The PAS, which includes accreta, increta, or percreta, can be causes of significant surgical and hemorrhagic morbidity on the labor and delivery floor\textsuperscript{4,37} (Figure 2). While PAS is relatively rare, particularly in the absence of a placenta previa, it can occur at vaginal delivery when there is no previa. Given

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\caption{The photo on the left (A) shows a retained portion of placenta approximately 8 weeks after delivery. The photo on the right (B) shows the same uterus following hysteroscopic morcellation of the retained placenta.}
\end{figure}

\begin{figure}[h]
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\includegraphics[width=\textwidth]{image}
\caption{Magnetic resonance image showing a portion of retained placenta 6 weeks postpartum. The arrow indicates an area where the light-gray placenta is deeply invasive into the darker-gray myometrium. Placenta accreta spectrum was confirmed pathologically following hysterectomy.}
\end{figure}
the excess morbidity, providers should consider this pathology when a placenta is retained in the setting of significant PAS risk factors. These include prior uterine surgeries, including hysteroscopic resections, IVF conception, a history of intrauterine adhesions, or a prior history of MAP or pathologic findings of accreta. 38,39

When a separation plane cannot be created or extraction attempts begin to invert the uterus, MAP should be suspected. In this case, further attempts to extract the placenta should cease, as forcible removal of a MAP can lead to massive hemorrhage. 40 At this point, consideration should be made for hysterectomy, which will be necessary if the patient has an undeliverable placenta with significant hemorrhage. Alternative treatment has been described including expectant management or uterine conservation. 41 Expectant management has been described in small studies and refers to the placenta left in situ after diagnosis of PAS. 38,41,42 Such management requires careful patient selection and counseling, as this risks delayed hemorrhage or infection. Nevertheless, successful conservative management has been described, with placental expulsion, resorption, or removal at a median of 3 months and up to 1 year postpartum. 43

Uterine conservation with placental removal is an alternative technique that likewise has been described in only small studies. This refers to resection of the placental bed at the area of suspected PAS and requires conversion to laparotomy after vaginal delivery. 41 The resultant defect in theory can be repaired via over-sewing and/or uterine repair or alternatively attempting tamponade with a Bakri balloon. Only one small study has evaluated the latter in a randomized control trial, and only with the lesser invasive types of PAS. 41,44

Conclusion
Retained placenta after vaginal delivery can be a source of significant hemorrhagic and surgical morbidity to the mother. In considering ways to lesson morbidity, the clinician should have a knowledge of risk factors for both retained placenta and MAP, allowing them to triage those patients most at risk of hemorrhage and prepare by ensuring blood products are easily available. When managing the patient with retained placenta, 30 minutes of elapsed third stage have been traditionally used as a guideline for timing manual removal; however, recent research has suggested that shorter duration of third stage may in fact be less morbid. Further research should be pursued to determine the best timing and infection prophylaxis for this etiology. Regardless, prompt diagnosis and management with appropriate personnel, access to blood for massive transfusion protocol, and surgical equipment such as uterine suction and tamponade can be required to treat retained placenta and lessen its morbidity.

Disclosure
The authors report no conflicts of interest in this work.

References

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