Abdominal Pregnancy in the Small Intestine Presenting as Acute Massive Lower Gastrointestinal Hemorrhage

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An abdominal pregnancy is an ectopic pregnancy in which the implantation site occurs in the abdominal cavity outside the female reproductive organs. There have been four reported cases that ruptured into the gastrointestinal tract and into the large intestine. We present the first case of an abdominal pregnancy rupturing into the small intestine with a good outcome.

1. Introduction

Acute lower gastrointestinal hemorrhage (LGIH) is defined as the onset of hematochezia, originating from either the colon or rectum. The most common causes of acute severe LGIH include diverticulosis, angioectasia, postpolypectomy bleeding, and ischemic colitis [1]. Ectopic pregnancy refers to the implantation of an embryo outside the uterus and is classified into two major types, according to the location of the implant—viz., a tubal or a nontubal pregnancy. The most common type of ectopic pregnancy (95%) is the tubal, which includes all parts of the fallopian tube (i.e., the fimbria, ampulla, isthmus, and cornual or interstitial part). An abdominal pregnancy is a nontubal pregnancy in which the implantation site occurs in the abdominal cavity and has a very low incidence (1%) [2]. Massive rectal bleeding is an unusual complication of ectopic pregnancy and carries with it a high mortality rate. We reviewed all cases of ectopic pregnancy that presented with LGIH [3–15] (Table 1) and only four of these were abdominal pregnancies.

For all reported cases of abdominal pregnancy presenting with lower GI bleeding, the gestational sac ruptured into the large intestine (colon or rectum). Herein, we present a case of an abdominal pregnancy in which the gestational sac ruptured into the small intestine and presented as severe hematochezia.

2. Case Presentation

A 32-year-old woman sought care at a provincial hospital after passing loose and dark stool about 10 times in a single day. She had been healthy until the diarrhea occurred and was not taking any medications. Her past medical history was unremarkable, except that she had undergone tubal surgery for pelvic inflammatory disease three years priorly. She had one child who was born by vaginal delivery. She had never undergone an instrumental pregnancy termination or intrauterine device insertion, which might lead to uterine perforation. She had no history of amenorrhea or abnormal vaginal discharge. Her initial diagnosis was upper gastrointestinal hemorrhage, but this was changed when bile was observed in the nasogastric tube after which the patient developed exsanguinating hematochezia and severe hypotension requiring 11 units of packed red cell transfusion for stabilization. Soon after, she was transferred to a tertiary care center.

The patient’s blood pressure was 70/40 mmHg, and her pulse rate was 120 beats/min. The abdominal examination was unremarkable. The per-rectal examination revealed excessive bleeding without any discernible cause. The laboratory tests showed a hemoglobin level of 6.3 g/dL and a platelet count of 28 x 10^3/μL. Owing to the unstable condition of the patient, an emergency exploratory laparotomy was conducted in order to localize and control the bleeding. During the laparotomy,
blood was found in the peritoneal cavity and a segment of the ileum attached to the fundus of uterus (Figure 1(a)). The intraluminal content was palpated in the adhered ileal segment (Figure 1(b)). An enterotomy revealed a fetus 7.5 cm in crown-rump length and fresh blood in the ileal lumen (Figure 1(c)). The placental tissue had implanted at the fundal dome of the uterus and eroded into the small bowel (Figure 1(d)). A segmental small bowel resection was performed along with reanastomosis. The placental tissue was removed by way of a wedge resection of the uterine wall. The patient had an uneventful postoperative course and was discharged on postoperative day 7.

Histologic examination of the resected specimens later confirmed the diagnosis of an abdominal pregnancy which included a male fetus of 4 months' gestational age (Figure 2), normal cord, subserous uterine myoma, and submucosal hemorrhage in the small intestine.

| Author                  | Reported year | Patient | Site of ectopic pregnancy | Location of placental erosion | Gestational age (weeks) | Maternal outcome | Fetal outcome |
|-------------------------|--------------|---------|---------------------------|-------------------------------|-------------------------|------------------|--------------|
| Armstrong [3]           | 1835         | NA      | NA                        | NA                            | 24                      | Dead             | NA           |
| Edgar [4]               | 1901         | NA      | NA                        | Sigmoid                       | NA                      | Dead             | NA           |
| Clark [5]               | 1932         | Female, 25 years | Left interstitial | Sigmoid                       | 6                       | Alive            | Dead         |
| Webster and Kerr [6]    | 1956         | NA      | Right interstitial        | Appendix and ileum            | NA                      | Dead             | NA           |
| Engel [7]               | 1961         | NA      | Left interstitial         | Ileum                         | NA                      | Dead             | NA           |
| Shirkey et al. [8]      | 1964         | Female, 38 years | Left interstitial | Ileum                         | NA                      | Alive            | Dead         |
| Bigg et al. [9]         | 1965         | Female, 19 years | Right interstitial | Caecum                        | NA                      | Alive            | Dead         |
| Bornman et al. [10]     | 1985         | Female, 29 years | Abdominal (posterior wall of uterus) | Sigmoid                       | 34                      | Alive            | Alive        |
| Seow et al. [11]        | 1992         | NA      | Abdominal                 | Caecum                        | NA                      | NA               | NA           |
| Verma et al. [12]       | 1996         | Female, 38 years | Left fallopian tube | Sigmoid                       | 4–8                     | Alive            | Dead         |
| Warshal et al. [13]     | 1996         | Female, 36 years | Right interstitial | Ileum                         | 12–16                   | Alive            | Dead         |
| Saravanane et al. [14]  | 1997         | Female, 30 years | Abdominal (pouch of Douglas) | Rectum                        | 14                      | Alive            | Dead         |
| Ekwaro et al. [15]      | 2004         | Female, 26 years | Abdominal (fundus of uterus) | Sigmoid                       | NA                      | Dead             | Dead         |
| Present study           | 2017         | Female, 32 years | Abdominal (fundus of uterus) | Ileum                         | 16                      | Alive            | Dead         |

**3. Discussion**

A case of acute massive lower gastrointestinal hemorrhage, caused by an abdominal pregnancy that ruptured into the small intestine, is discussed. This is a rare gynecologic condition, considered to be an underlying cause of rectal bleeding, and is usually misdiagnosed, resulting in delayed treatment. Due to the rarity of the condition and the associated high mortality of abdominal pregnancy, despite having a high index of suspicion, it is not possible to diagnose the condition before surgery.

Lower gastrointestinal hemorrhage is defined as bleeding originating distal to the ligament of Treitz. Typically, massive bleeding is thought to require more than 3 to 5 units of blood transfused over 24 h. Although LGIH can occur at any age, the disease presentation for adults tends to be diverticular bleeding, inflammatory bowel disease, or neoplasm, and with advancing age, bleeding from an arteriovenous malformation, diverticular bleeding, or neoplasm [16]. The other common causes of LGIH are ischemic colitis, postpolypectomy bleeding, hemorrhoids, and stercoral ulcer.

We searched for reports on intestinal hemorrhage associated with ectopic pregnancy and found only 13 cases (Table 1). The site of the ectopic pregnancy was not identified in the first two reported cases. The first [3] was a case of a syphilitic woman six months pregnant, dying of hemorrhage and suddenly passing bloody stool containing fetal bones, and the second [4] was a case of rectal hemorrhage, proven at autopsy to have arisen from an ectopic gestational sac rupture into the sigmoid colon. The other 11 reports in the literature identified the implantation sites; for most of which (six cases) the implantation site was at the uterine serosa and typically ruptured into the large intestine. The remaining case was a tubal ectopic pregnancy that ruptured into the sigmoid colon.
All types of ectopic pregnancies result in high maternal mortality, caused by delayed treatment of life-threatening hemorrhages. It is estimated that a woman with an abdominal pregnancy is 90 times more likely to die compared to a woman with an intrauterine pregnancy [17]. Historically, lack of knowledge of the condition was associated with mortality of the mother. Even today, the fetus almost always dies in ectopic pregnancies, although we did find one case of an abdominal pregnancy that presented as massive rectal bleeding, where both the mother and neonate survived [10].

The abnormal implantation site of an ectopic pregnancy leads to denigration of the trophoblastic tissue [13], which grows by invading the adjacent structures that are prone to the implantation site where there is greater blood supply. The communication of the intestinal tract with the ectopic gestational sac is called an enteroamniotic fistula, which follows the villous invasion of the bowel wall. When the gestational sac approximates to the intestine, there is an inflammatory reaction and an infection develops, resulting in a fistula. The vascularized gestational structures are aggravated by infection and villous invasion of the adjacent vascular structures, leading to massive hemorrhage [8]. The terminal ileum, sigmoid colon, and caecum are the parts of the gastrointestinal tract that are usually involved [14]. The small intestine is associated with interstitial and not abdominal pregnancies, which are usually associated with invasion of the large intestine. We have thus presented the first case of a ruptured abdominal pregnancy into the small intestine, which caused massive lower gastrointestinal hemorrhage.

Attending physicians need to be aware of the possibility of a fistula between the ectopic gestational sac and the bowel in any pregnant woman who presents with obscure intestinal hemorrhage, and to initiate prompt surgery. Although women with ectopic pregnancy frequently have no identifiable risk factors, a prospective case-controlled study revealed that increased awareness of ectopic pregnancy and knowledge of the associated risk factors help to identify women at higher risk and early diagnosis. In our case, we did not consider this condition initially but performed an emergency surgery after massive blood transfusions. During laparotomy—in order to control the bleeding and to prevent intraabdominal abscess formation or peritoneal sepsis—we resected the infected fistula, a segment of the small bowel, the wall of the involved uterus, and the ectopic fetus and placental tissue. After recovering from the surgery, the patient was discharged without any major morbidity.

![Figure 1: Photographs of intraoperative findings. (a) Exploratory laparotomy showing the adherence between the ileum and uterine fundus. (b) The intraluminal ileal mass palpable near the adherence site. (c) The opened ileum revealing a dead fetus. (d) Photograph showing the placenta in the uterus after detaching the ileum from the uterine fundus.](image1)

![Figure 2: Photograph of the dead fetus.](image2)
In conclusion, abdominal pregnancy is a rare life-threatening condition requiring prompt treatment. This is the first report of an abdominal pregnancy that ruptured into the small intestine with a good outcome.

Ethical Approval
This study was approved by the Institutional Review Board, IRB (HE601262), Khon Kaen University.

Conflicts of Interest
The authors had no conflicts of interest regarding publication of this paper.

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