CASE REPORT

Case report: successful open resection of a symptomatic giant liver haemangioma during the second trimester of pregnancy

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Abstract

Liver haemangiomas are consistently reported to be the commonest benign liver tumours and are most often incidental findings in asymptomatic individuals. Large lesions can become symptomatic, spontaneously rupture or result in a consumptive coagulopathy known as Kasabach–Merrit Syndrome. It is believed that the female sex hormones, particularly oestrogen, have an influence on haemangioma development and growth. The optimal management of large haemangiomas during the hyperoestrogenic state of pregnancy is poorly described in the current literature. To our knowledge, we describe only the second case of a giant hepatic haemangioma resection during pregnancy.

INTRODUCTION

Haemangiomas are the most common benign liver neoplasm and result from ectatic growth of the vasculature [1]. These lesions can grow to a significant size and are referred to as giant haemangiomas once larger than 5 cm. Despite the absence of invasive or metastatic potential, hepatic haemangiomas can result in pain, obstructive symptoms, rupture with life threatening haemoperitoneum or a consumptive coagulopathy known as Kasabach–Merrit syndrome [2]. Consensus agreement on the indications for surgical resection, other than for symptomatic lesions, including for giant haemangioma, is lacking at present [1]. There is a paucity of available literature outlining the behaviour of haemangiomas during pregnancy, however female sex hormones have been implicated in their pathogenesis and growth [3]. The nature and extent of this relationship is yet to be fully elucidated. On the other hand, the diagnosis of a symptomatic giant haemangioma during pregnancy presents a dilemma because the safety of neither conservative nor surgical resection has been rigorously investigated. At our centre, a 30-year-old G2P0M1 female underwent resection of a symptomatic giant hepatic haemangioma during the second trimester without any complication.

CASE REPORT

An otherwise well 30-year-old female presented to her general practitioner complaining of intermittent right upper quadrant discomfort over the preceding months. Abdominal examination
was suspicious for hepatomegaly and subsequent ultrasound demonstrated a large liver mass. The patient was referred to hospital and CT imaging confirmed a pedunculated giant hepatic haemangioma (13.8 cm x 10.2 cm x 7.4 cm) attached to segment 4 of the liver (Figs 1 and 2). Two smaller intraparenchymal haemangiomas in segment 6 and 8 were also identified and measured 0.5 cm and 1.3 cm, respectively. Despite a negative urine Beta HCG test at the time of CT scanning, she was subsequently found to be pregnant. False negative results such as this are reported to occur, especially in the early luteal phase \[4\] when the urine HCG may not have reached the diagnostic threshold. Approximately 1 year prior, she had ceased the combined oral contraceptive after taking it for 10 years and was planning to conceive.

On first review in our clinic the patient was 6 weeks pregnant. She was counselled by a consultant liver surgeon and obstetrician about the available management options and associated risks. The surgical options discussed included; close observation throughout pregnancy and elective resection postpartum, termination of pregnancy followed by resection or resection during the second trimester of pregnancy. Our patient decided to have the tumour resected whilst pregnant due to the concern of rupture, accepting the risks of surgery to herself and the foetus. The operation was scheduled for the second trimester as this is the period in which subsequent miscarriage or preterm labour are least likely \[5\].

The resection was performed at 18 weeks gestation under general anaesthesia via a small upper midline incision. Upon entering the peritoneal cavity the exophytic haemangioma (Fig. 3) was resected along its attachment to segment 4b using Harmonic ACE®+7 Shears. This allowed removal with minimal disruption of liver parenchyma, major vasculature or bile ducts. Minimal blood loss occurred and operating time was approximately 1 h. The smaller intra-parenchymal lesions were not resected as they were not the likely cause of her symptoms and represented a lower risk of rupture. The general anaesthetic was tailored to reduce any interruptions to uterine blood flow. Intraoperative foetal monitoring was not indicated due to the pregnancy still being in the previable phase.

Obstetric examination in the immediate post-operative period was consistent with a healthy foetus and the patient was discharged from hospital on the third post-operative day. Histopathological examination of the tumour confirmed a haemangioma weighing 412 g. The patient received outpatient follow-up and did not experience any surgical or obstetric complications. A healthy infant was delivered at term via an elective caesarean section due to maternal request.

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Figure 1: Coronal CT image in the portal venous phase demonstrating the large exophytic haemangioma.

Figure 2: Axial CT image in portal venous phase at level slightly inferior to aortic bifurcation demonstrating the peripheral nodular enhancement of the haemangioma.

Figure 3: Intraoperative image demonstrating the exophytic nature of the haemangioma and relevant anatomical structures.
DISCUSSION

There is little available literature describing the outcome of liver haemangioma resection during pregnancy. To our knowledge, the only available publication is an isolated case report of a successfully resected giant haemangioma at 16 weeks gestation due to Kasabach–Merit Syndrome [6]. The scarcity of data may be representative of how infrequently this operation is performed during pregnancy. Both pregnancy and hormonal therapy, including the oral contraceptive, have been associated with haemangiomas enlargement [3]. Descriptions of tumour growth and complications during pregnancy are conflicting. Rapid growth [7] and haemorrhagic complications [8] have been reported but a recent small case series (four patients, seven pregnancies) found that these tumours can be safely observed during pregnancy [2]. A larger observational cohort study in non-pregnant individuals found that haemangioma resection can be justified due to minimal morbidity and no mortality in the 13 patients who underwent surgery [9]. However, these authors also found that no tumours under 5 cm increased in size or ruptured in patients who received conservative management. Obviously, extrapolation of these findings to our patient would be misguided as the large peripheral exophytic tumour was >5 cm and she was pregnant.

Spontaneous rupture is the complication most feared because this is associated with a mortality rate approaching 80% [6]. A recent large study has estimated that the rupture rate of giant liver haemangioma is 3.4% with peripherally located and exophytic lesions being at higher risk [10]. Accelerated growth, increased intra-abdominal pressure and direct contact with the gravid uterus are all plausible mechanisms for spontaneous rupture or worsening symptoms during pregnancy. Intrapartum rupture of liver haemangiomas has been reported, supporting the notion that increased intra-abdominal pressure may be a causative factor. Our patient’s tumour was assessed as being high risk for complications during pregnancy due to its size, exophytic nature and subumbilical position. Therefore, we felt that the risks associated with surgery were justified in this case as spontaneous or intrapartum rupture could be an unsurvivable event.

To our knowledge, this is only the second case report of a successful hepatic haemangioma resection during pregnancy. The natural history of these lesions during pregnancy is incompletely understood and resection should be considered if worsening symptoms or the clinical concern for rupture is significant. An interdisciplinary approach that involves surgical, anaesthetic and obstetric specialists is required to perform this procedure safely, and a good overall result for mother and child is possible.

CONFLICTS OF INTEREST STATEMENT

None to declare.

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