

CASE REPORT

The entrapped twin: a case of fetus-in-fetu

Rashide Yaacob, Annisa Zainal Mokhtar, Dyg Zahratul Hamrak Abang Jamari, Norliana Jaafar

Department of General Surgery, Sultan Abdul Halim Hospital, Sungai Petani, Malaysia

Correspondence toDr Rashide Yaacob,
edie14770@yahoo.com

Accepted 24 August 2017

SUMMARY

Fetus-in-fetu (FIF) is a rare entity in which malformed parasitic twin grows inside the body of its twin. It is most commonly presented with mass in the abdomen. We present a case of a 15-year-old boy who presented with abdominal mass since infancy. Radiological investigations are suggestive of FIF. Intraoperatively, malformed fetus in a sac was found and excised. Postoperatively the patient recovers well and was put on follow-up.

BACKGROUND

Fetus-in-fetu (FIF) is an uncommon pathology with the incidence of 1 per 500 000 births. There are many theories proposed with regard to the aetiology of FIF. One of the theories which is increasingly proposed in the recent literature is that the FIF is a diamniotic, monochorionic, monozygotic twin that becomes incorporated into the body of the host twin after anastomosis of the vitelline circulation.² Other theories include demised multiple pregnancy³ and variant of mature teratoma. 4 Most common presentation is a mass in the abdomen.⁵ Other sites include intracranial, mediastinum, sacrum, scrotum and mouth.6 It can also present with a mass effect. Diagnosis can be made via X-ray, ultrasound, CT scan and MRI. Some cases were diagnosed after prenatal screening.⁵ Another diagnostic modality (molecular analysis) is using an informative genetic marker for uniparental isodisomy of chromosomes 14 and 15. If there is no genetic difference between the host and the fetiform mass, then it is diagnostic of FIF.⁵ FIF has to be distinguished from teratoma as the latter has the potential to recur though FIF with malignant recurrence has been reported. Complete excision of the mass is curative and will help in ruling out teratoma with malignant elements.

CASE PRESENTATION

A 15-year-old boy presented to our hospital with history of abdominal swelling since birth which progressively enlarged recently. He also complaints of abdominal pain and was unable to tolerate orally for 1 week prior to admission. On examination, there was tender, hard mass over central part of abdomen.

INVESTIGATIONS

CT abdomen and pelvis revealed a huge intra-abdominal mass $14 \times 18.5 \times 23.8$ cm that extends up to the subhepatic region with inferior extension into anterior pelvis. Components in favour of a fetus

that were seen within the mass include deformed skull, vertebral body and long bones (figure 1). CT angiography was done to localise the feeding vessels involved. CT angiography shows multiple small arteries supplying the wall of the mass. No main arterial pedicle detected (figure 2). No tumour markers were taken preoperatively.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of this condition would be organised teratoma.

TREATMENT

Laparotomy was performed through a midline laparotomy incision with findings of a 15×10 cm intraperitoneal mass with multiple huge feeding vessels densely attached to small bowel mesentery. The mass was perforated over pelvic region with 1000 cc pus aspirated. A gestational sac weighing 2.5 kg was delivered, and when opened, a 1.6 kg non-viable baby noted with shortened and malformed upper and lower limbs, hypoplastic trunk, long hair, developed male genitalia with pubic hair, imperforated anus, fused malformed eyes, vertebra and normal baby skin covered with vernix caecosa. There was no mouth, umbilical cord or placenta (figure 3).

OUTCOME AND FOLLOW-UP

Postoperatively, the baby was returned to the family for ritual funeral as requested by family. Histopathological examination (HPE) of the sac enveloping the fetus revealed a fibrous cyst wall with focal area lining of mature squamous epithelium with many hair shafts and some muscle in subepithelial stroma. The postoperative period of the patient was uneventful. Alpha-feto protein (AFP) taken 3 months postoperation was normal and the patient has no active complaints during subsequent follow ups.

DISCUSSION

It is still a controversy as to whether FIF is an entity on its own or is a highly organised teratoma. Teratoma is a true neoplasm composed of multiple tissues foreign to the part in which it arises.² Some writers emphasise the presence of vertebral column as distinct feature in diagnosing FIF.⁵ However, there are reports of FIF with differentiation of tissues without vertebral column.⁸ In our case, although only HPE of the sac was obtained as the baby was returned to family members after the operation for funeral according to local custom,



To cite: Yaacob R, Zainal Mokhtar A, Abang Jamari DZH, et al. BMJ Case Rep Published Online First: [please include Day Month Year]. doi:10.1136/ bcr-2017-220801



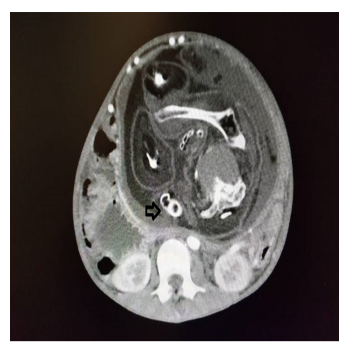


Figure 1 CT abdomen showing an intra-abdominal mass with components in favour of a fetus within the mass, including deformed skull, vertebra body and long bones. Arrow shows vertebra body.

the diagnosis of FIF was confirmed via the presence of vertebral columns in CT abdomen.

Treatment of FIF is surgical. Although benign, complete excision of FIF together with the capsule is crucial as there is a possibility of malignant recurrence if any of the tissue is not completely excised. Surgical excision also allows for relieve of obstruction and prevention from further compression.

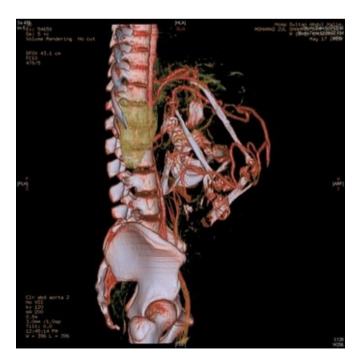


Figure 2 CT angiography of the abdomen and the pelvis showing multiple small arterial supplying the wall. No main arterial pedicle detected.



Figure 3 Malformed fetus with hypoplastic trunk, long hair, developed male genitalia, fused eyes and normal baby skin.

The operation to remove FIF is a challenging operation as the mass is highly vascular with multiple feeding vessels. The big size of the FIF has made surgery more difficult as there is high possibility of injuring surrounding structures. CT angiography can help in identifying obvious vascularity and help in assessing the feasibility of doing such cases in the involved centre. In our case, CT angiography was done as this is a rare case and the first one encountered in our centre. With limited experience in handling such cases, CT angiography has given a rough idea on the possible difficulties that may be encountered by us intraoperatively especially with regards to vascular supply of the mass.

Learning points

- ► Mass per abdomen with axial skeleton or long bone seen during radiological must raise suspicion of fetus-in-fetu (FIF).
- Operation to remove the mass must involve careful planning as there is high possibility of bleeding intraoperatively as the mass is highly vascularised. This involved CT angiography to review the feeding vessel.
- ► FIF must be differentiated from teratoma as there is a risk of recurrence in teratoma.
- ► Further follow up is needed as although the mass is proven to be FIF, there is still a small risk of malignant recurrence especially if there is inadequate removal of sac during operation.

Contributors RY: gave the idea for the article; guarantor. AZM, DZHAJ and NJ: performed literature review. All authors: wrote the article and identified and managed the case.

Competing interests None declared.

Patient consent Guardian consent obtained.

Provenance and peer review Not commissioned; externally peer reviewed.

Open Access This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work

is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

© BMJ Publishing Group Ltd (unless otherwise stated in the text of the article) 2017. All rights reserved. No commercial use is permitted unless otherwise expressly granted.

REFERENCES

- 1 Grant P, Pearn JH. Foetus-in-foetu. Med J Aust 1969;1:1016-20.
- 2 Willis RA. The borderland of embryology and pathology. 2nd edn. Washington, DC: Butterworths, 1962:442–62.
- 3 Pang KK, Chao NS, Tsang TK, et al. From observation to aetiology: a case report of a twin fetus-in-fetu and a revisit of the known rarity. Hong Kong Med J 2015;21:80–3.

- 4 Hopkins KL, Dickson PK, Ball TI, et al. Fetus-in-fetu with malignant recurrence. J Pediatr Surg 1997;32:1476–9.
- 5 Brand A, Alves MC, Saraiva C, et al. Fetus in fetu-diagnostic criteria and differential diagnosis-a case report and literature review. J Pediatr Surg 2004;39:616–8.
- 6 Abdulraheem NT, Nasir AA, Abdur-Rahman LO, et al. Oral fetus-in-fetu: a case report. J Pediatr Surg Case Rep 2015;3:171–3.
- 7 Spencer R. Parasitic conjoined twins: external, internal (fetuses in fetu and teratomas), and detached (acardiacs). *Clin Anat* 2001;14:428–44.
- 8 Mohta A, Shrivastava UK, Sodhi P, et al. Fetus-in-fetu. Pediatr Surg Int 2003:19:499–500.
- 9 Ji Y, Song B, Chen S, et al. Fetus in fetu in the scrotal sac: case report and literature review. Medicine 2015;94:e1322.
- 10 Lindsey M, William J, Tyler A. Fetus in fetu: review of the literature over the past 15 years. J Ped Surg Case Reports 2015;3:554–62.

Copyright 2017 BMJ Publishing Group. All rights reserved. For permission to reuse any of this content visit http://group.bmj.com/group/rights-licensing/permissions.

BMJ Case Report Fellows may re-use this article for personal use and teaching without any further permission.

Become a Fellow of BMJ Case Reports today and you can:

- ► Submit as many cases as you like
- ► Enjoy fast sympathetic peer review and rapid publication of accepted articles
- Access all the published articles
- ▶ Re-use any of the published material for personal use and teaching without further permission

For information on Institutional Fellowships contact consortiasales@bmjgroup.com

Visit casereports.bmj.com for more articles like this and to become a Fellow