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Davek Sharma
*Abington Jefferson Health*

Jacob Katsnelson
*Abington Jefferson Health*

Emmanuel Nwachuku
*Abington Jefferson Health*

Jeffrey Kolff
*Abington Jefferson Health*

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The De Garengeot hernia: A case report of an unusual presentation of appendicitis

Davek Sharma *, Jacob Katsnelson, Emmanuel Nwachuku, Jeffrey Kolff

Department of General Surgery, Abington Hospital, Jefferson Health, 1200 Old York Rd., Abington, PA, 19001, United States

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ABSTRACT

INTRODUCTION: A De Garengeot hernia is defined by a femoral hernia containing the appendix. Acute appendicitis within a femoral hernia is an extremely rare surgical presentation and occurs in only 0.08–0.013% of cases as cited by the literature.

PRESENTATION OF CASE: A 64-year-old female presented to the Emergency Department of our hospital with acute onset of a right-sided groin bulge that occurred earlier that day after doing heavy lifting. Her workup revealed acute appendicitis contained within an incarcerated right femoral hernia. The patient underwent laparoscopic appendectomy with open femoral hernia repair. Intraoperatively, the appendiceal tip was incarcerated within the hernia sac. It was removed through the open inguinal incision after the appendix base was divided laparoscopically. Final pathology showed inflamed acute appendicitis without evidence for neoplasm.

DISCUSSION: Physicians should be aware of the rare entity of an unusual presentation of appendicitis as well as surgical options for treatment. The literature does not conclude upon a gold standard for method of approach.

CONCLUSION: De Garengeot hernia remains a rare and unusual surgical presentation of femoral hernia, and complication of the case by incarceration leading to acute appendicitis provides a challenging surgical approach which should be individualized to each patient.

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1. Introduction

Femoral hernias occur as a weakness and protrusion of abdominal viscera or contents through the femoral canal, bordered superiorly by the inguinal ligament and laterally by the femoral vessels. The femoral hernia is much more common in women and accounts for 3% of all hernias [1]. Due to its narrow neck, it has a much larger chance of strangulation compared to inguinal hernias and all are recommended for repair [1,2]. The migration of the appendix into the inguinal hernia sac is known as an Aymard hernia and is very rare. However, even more rare is the migration of the appendix into the femoral hernia sac, which is called De Garengeot hernia, with a reported incidence of 0.5–5% of all femoral hernias [2]. The incidence of De Garengeot's hernia cited in the literature is fewer than 100 cases [2]. The incidence of concurrent appendicitis in the presence of a De Garengeot hernia is cited to be as low as 0.08%–0.13% [3]. We present a rare case of an acutely inflamed appendicitis within a De Garengeot hernia diagnosed preoperatively and managed surgically.

2. Case

This case report has been written in accordance with the SCARE 2018 criteria [4].

A 64-year-old female presented to the Emergency Department of our hospital with acute onset of a right-sided groin bulge that occurred earlier that day after doing heavy lifting. She had no significant past medical or surgical history. Her baseline functional status was exceedingly well. Her laboratory studies showed no leukocytosis or any electrolyte abnormalities. Initial radiologic studies included a transabdominal and transvaginal pelvic ultrasound, which showed a 2.4 × 2.6 cm fluid collection with some septations and absent visualized adnexal structures. A subsequent CT demonstrated a well-circumscribed 2.7 × 3.2 × 2.6 cm fluid collection containing echogenic debris. It did not have significant rim enhancement yet was noted to be most consistent with an abscess, complex seroma, or necrotic lymph node; its depth was amenable to percutaneous drainage. On further review with the radiologist, it was noted to contain the appendix, the tip of which was dilated and radiographically concerning for a cystic process, with mucocoele remaining within the differential diagnosis.
3. Discussion

The De Garengeot hernia was first described in 1731 by French surgeon Rene Jacques de Garengeot [3]. The leading theories behind the presence of appendix within the femoral canal emphasize an embryologic abnormal pelvic appendiceal location on the cecum, predisposing it to femoral canal herniation [3,5], and a larger cecum which causes the appendix to herniate by mass effect [3]. The presence of appendicitis within the De Garengeot hernia is also debated with regards to the sequence [5]. Most seem to favor a theory of appendicitis resulting from extrinsic compression of the appendiceal lumen by a narrow femoral neck, as opposed to appendicitis happening first and then migrating down into the femoral hernia [3].

Similar to inguinal hernia, De Garengeot hernia clinically presents with a groin lump, which is nonreducible and tender to palpation [2–4]. Fever or bowel obstruction are possible but rare (Figs. 3 and 4).

The diagnosis is usually made preoperatively with modern imaging techniques, but there are reported cases in the literature of diagnosis being made intraoperatively [6]. Treatment is a surgical emergency. There is no specific agreed upon surgical procedure for management [6,7]. For a noninflamed appendix, laparoscopic appendectomy with intraperitoneal mesh repair might be a consideration. Beysens et al. reported the first case of laparoscopic appendectomy followed by TEP femoral hernia repair [8]. We recommend avoidance of mesh-based repair techniques in the setting of appendicitis, specifically perforated cases. Advocates of pure laparoscopic approaches support potential advantage in a shorter hospital stay and speedier recovery [9]. The most common complications are wound infections [5].

De Garengeot hernias are rare, and a concomitant diagnosis of acute appendicitis within the hernia is exceedingly rare. We believe the rarity of this case can contribute to the surgical literature to provide our institutional experience and allow practitioners to be prepared with their own thoughts for management should they ever encounter this rare surgical entity.
4. Conclusion

De Garengeot hernia remains a rare and unusual surgical presentation of femoral hernia, and complication of the case by incarceration leading to acute appendicitis provides a challenging surgical approach which should be individualized to each patient. As there are no guidelines for surgical management for this exceedingly rare entity, knowledge of the condition and reporting of presentations and treatments provides the surgical community with knowledge for potential approaches.

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Ethical approval

We did not seek IRB approval for this case report, which contains only retrospective, deidentified patient information. The writing or publication of this case report did not affect this patient’s treatment or outcomes in any way. There are no ethical dilemmas with this case.

Consent

Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

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Davek Sharma: Conceptualization, Methodology. Writing - original draft. Jacob Katsnelson: Investigation, Methodology. Emmanuel Nwachuku: Writing - review & editing. Jeffrey Kolff: Supervision.

Declaration of Competing Interest

The authors report no declarations of interest.

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