Ileal duplication cyst in the elderly complicated by appendicitis: A rare case report and review of literature

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1. Introduction

Gastrointestinal duplication cysts are rare and generally manifest in the pediatric age group with acute abdomen. The entity is rare in adults with only few cases reported so far. They may arise anywhere in the alimentary tract from the mouth to the anus. They are connected to the bowel with a well-defined muscular coat and characteristically lined by gastrointestinal mucosa. Adults are generally asymptomatic or may have vague abdominal symptoms [1]. Obstruction, hemorrhage, perforation and malignancy may complicate the condition. We present here one such rare case of terminal ileal duplication cyst with reactive appendicitis in an elderly lady that was managed laparoscopically.

2. Presentation of the case

A 61 years lady presented with generalized abdominal pain, on and off since 6 months that was conserved with oral medications (antibiotics and analgesics) but had increased in severity in the last two months. There was no associated vomiting, fever or change in bowel or bladder habits. Her physical examination was unremarkable except for mild tenderness in the right iliac fossa. She had an elevated leukocyte count (14,000/cu/mm3). Ultrasonography (USG) of the abdomen done at another institute showed a cystic structure in the right lower abdomen. We further evaluated her with a Computed Tomography (CT) that revealed a well-defined cystic bilobed lesion (7.6 × 4.7 × 3.5 cm) extending from the intramural portion of the medial wall of the caecum to immediately posterior to the ileocaecal junction with multiple enlarged ileocaecal mesenteric lymph nodes (Fig. 1A & 1B). The Appendix was seen separately. The findings were suggestive of an ileocaecal duplication cyst with secondary infection. Intraoperatively, a mass was found just proximal to the ileocaecal junction adherent to the surrounding mesentery with multiple enlarged mesenteric lymph nodes. A laparoscopic right hemicolectomy with side to side stapled anastomosis was done. Grossly on cut opening the specimen, a bulging cyst at the terminal ileum measuring 5.5 × 5 cm, filled with clear mucoid fluid. It had a flat lining with a focally granular appearance (Fig. 2). Histopathology confirmed a terminal ileal duplication cyst (Fig. 3) with reactive appendicitis and reactive follicular hyperplasia in the involved lymph nodes. The cyst was non-communicating with the lumen. The cyst lining was found to be columnar with villiform projections lined by columnar foveolar type gastric epithelium but no inflammation on high and low resolution (Fig. 4A&B). No malignant degeneration was seen. The post-operative period was uneventful and the patient was discharged after five days. The patient is on regular follow-up and symptom free.

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Fig. 1. A Contrast Enhanced CT scan (coronal view) showing a well-defined bilobed cystic lesion (7.6 × 4.7 × 3.5 cm) involving ileocaecal junction. B Contrast Enhanced CT scan (axial view) showing the cystic lesion non-communicating with the lumen.

3. Discussion

Duplication cyst is a rare congenital anomaly presenting in pediatric patients that is most commonly seen in the ileum [2]. It was first described by Wendel [3]. There have been few cases of its presentation in adults reported ever since. The condition is usually seen in males. A number of etiological theories have been suggested, of which persistence of the fetal gut diverticula or ‘aborted Gemini’, and defect in the recanalization of the primitive gut are the most accepted. Any failure in the transition process of the intestinal tract from solid to tubular state between 6 and 8 weeks of intrauterine life may lead to the formation of duplication cysts [4]. These cysts have an outer muscular lining with an inner gastrointestinal

Fig. 2. Gross specimen showing collapsed cyst (non communicating) in the lumen of terminal ileum, Appendix seen.

Fig. 3. A Microphotograph of the duplication cyst H & E stain, 40 x magnification, showing the ileal mucosa below and the gastric mucosa showing foveolar epithelium with mucous glands beneath. A muscle wall is seen in between which is partially fibrosed. B Microphotograph of the Appendix showing the reactive follicles suggesting appendicitis.
Table 1
Overview of reported duplication cysts in adults.

| S. No. | Author                  | Age/Sex and relevant findings.                                                                 | Diagnosis                                                                                     | Lap/open                  |
|--------|-------------------------|-----------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------|
| 1      | Collaud S et al. [12].  | Adult, right lower abdomen pain. Pre-op imaging showed infected cyst.                         | Completely isolate ileal duplication cyst with low grade mucinous cystadenoma.                  | –                        |
| 2      | Isamu Hoshino et al. [13]. | Adult, abdominal pain. CT showed cystic mass in the small bowel with intussusception.    | Ileal duplication cyst                                                                         | Laparotomy with small bowel resection and anastomosis.                         |
| 3      | Tomas et al. [14].      | Adult, asymptomatic with past history of bilateral breast cancer. CT: non communicating cystic lesion in the terminal ileum | Mucinous cystadenoma in isolated ileal duplication cyst (on antimesenteric side)               | Surgical resection.          |
| 4      | Otter M.I. et al. [15]. | Adult, 52yrs/M, urinary symptoms.                                                             | Intestinal duplication cyst.                                                                  | Surgical resection.          |
| 5      | Smith J.H. et al. [16].  | Adult                                                                                       | Ileal duplication cyst with carcinoma                                                          | Surgical resection.          |
| 6      | Kim H.S. et al. [17].   | 19yrs/F, rt. Upper quadrant pain, CT showing ileocolic intussusception with cystic mass at tip. | Ileal duplication cyst                                                                         | Surgical resection.          |
| 7      | Gregor Blank et al. [18]. | 51yrs/M, asymptomatic.                                                                      | Ileal duplication cyst with adenocarcinoma.                                                   | Surgical resection of cyst without bowel resection.                            |

Fig. 4. A 400 x resolution histopathology slide showing cyst wall lined by columnar foveolar type gastric epithelium and at places thrown into villiform projections. No goblet cells are seen. Sub-epithelium shows mucous glands. B 100 x resolution showing gastric mucous lining characterized by columnar foveolar type gastric epithelium. Sub-epithelium shows multiple mucous glands.

In adults, the condition may be left undiagnosed. Commonly, they present with vague abdominal complaints. However, they may also present acutely in the setting of obstruction, hemorrhage, or perforation [5]. Due to the risk of progression from benign pathology to malignancy or other complications, surgery should be offered, preferably minimally invasive approach to all asymptomatic or asymptomatic patients. Rarely, the cyst may cause a fistula with the adjoining structure [6]. Duplication cysts may be associated with malrotation, esophageal duplication and vertebral anomalies [3, 7]. A cyst may be termed as a true duplication cyst if it fulfills the three criteria: attachment to any part of the gastrointestinal tract, lined by any gastrointestinal epithelium and containing smooth muscle in its wall [8]. It may be cystic or tubular. Cystic duplication cysts are commoner. They do not communicate with the bowel, but may share a common wall and blood supply. Tubular duplication cysts communicate with the intestinal lumen [8].

The preoperative diagnosis can be made on USG and CT. The cystic nature of the lesion is visible on USG. The diagnosis can be confirmed even on USG if the characteristic inner echogenic layer and the hypoechoegenic outer muscular layer can be defined [2]. Kumar et al. proposed a five layered cyst wall sign seen on ultrasonography delineating the outermost hyperechoic serosa, hypoechoic muscularis propria, hyperechoic submucosa, hypoechoic muscularis mucosa and the inner most hyperechoic mucosa [9]. CT scan shows the fluid filled cystic structures or the tubular structures with slightly enhancing walls in close proximity to the small bowel. Endoscopic ultrasound may also be used for the evaluation of duplication cysts. Surgery is imperative in symptomatic patients. There is no data available for the management in asymptomatic patients. However, it is deemed beneficial to surgically treat these cases due to the possible malignant change in duplication cysts [10]. Surgery entails resecting the involved bowel, however, in completely isolated cysts; excision of these may be done without resecting the bowel. Ileal duplication cysts are known to show heterotopic gastric mucosa [11].

Amongst the adult cases presented so far, some had degenerative changes in the cyst leading to malignancy. Table 1 gives a brief overview of the same. Our case did not show any such degenerative changes.

4. Conclusion

We highlight the rare occurrence of terminal ileal duplication cyst in the elderly. Surgical treatment is necessary due to the probable malignant transformation of duplication cysts. Laparoscopy adds on the advantage of minimally invasive approaches. Though rare, a differential diagnosis of duplication cyst should be con-
sidered in adult patients with cystic lesions of or in proximity to gastro-intestinal tract.

Conflict of interest

None.

Funding

None.

Ethical approval

Ethics Committee, Lilavati Hospital and Research Center.

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.

Author contribution

Kamal Yadav and Priyanka Sali wrote the paper.
Bhushan Bhole analysed the data.
Hitesh Mehta and Chandralekha Tampi performed the study.

Guarantor

Kamal Yadav.
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