CASE REPORT

Double appendixes: A rare case report

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Abstract

Anomalies of the appendix are rare, and one of the rarest is the double appendixes. Most anomalies of the appendix are observed in adults and are discovered incidentally during surgery that does not primarily involve the appendix. It is usually missed, often with life-threatening consequences.

KEYWORDS

anatomical variations of vermiform appendix, appendiceal duplication, appendicitis, double appendixes

1 | INTRODUCTION

Anomalies of the appendix are rare, and one of the rarest is the double appendixes. Most anomalies of the appendix are observed in adults and are discovered incidentally during surgery that does not primarily involve the appendix. There are many types of double appendixes, with some of the cases including duplication of other organs. It is usually missed unless a person undergoes a surgery that reveals it, often with life-threatening consequences. Herein, we present a rare case of a patient with double appendixes having acute appendicitis of both appendixes.

2 | CASE REPORT

A 30-year-old female patient with a height of 1.62 m and weight of 110 kg (BMI 41.91) having pain in the lower abdomen was admitted to the emergency department. The previous day, symptoms started with pain in the epigastrium. As time went on, the pain was located around the navel and finally settled in the right iliac fossa. The patient did not report having nausea, episodes of vomiting, or fever at home. The patient’s bowel habits were unaltered, and she reported loss of appetite the previous two days.

The physical examination raised the suspicion of appendicitis since a positive McBurney’s sign was found—right
lower abdominal quadrant pain—and rebound tenderness, indicating peritoneum irritation. The blood test results revealed leukocytosis, with WBC 12.730/mm³ (66.4% Neut, 23.6% Lym, and 8.1% Mono) and increased levels of the C-reactive protein (2.06). Our patient had an Alvarado score 9.³ Additionally, neither the urine analysis nor the abdominal X-ray revealed any findings.

Afterward, an examination of the abdomen was performed with an ultrasound. The coarse examination revealed no particular findings. The graded compression technique was then performed using the 8 MHz probe set above the position of maximum sensitivity with gradually increasing pressure exerted to displace the normal supernatant gas. The appendix was identified (Figure 1) with the blind end of the appendix arising from the base of the cecum. The appendix was dilated (diameter 1.5cm) in target appearance (axial section) (a) was non-compressible when compression was applied (b) and had hyperechoic appendicolith with posterior acoustic shadowing and periappendiceal fluid collection (white arrow) (c). After the application of the colored Doppler, it showed intense vascularization of the wall as an image of mural hyperemia (d).

The patient was hemodynamically stable but, during admittance, presented with a low-grade fever (37.5°C). After reviewing the test results, a decision was made to perform an open appendectomy via a McBurney’s incision.

Intraoperative findings included a mild quantity of free fluid in the right iliac fossa and an inflamed appendix. Surprisingly, another thin, mildly inflamed appendix was found when the appendectomy was completed (Figure 2).

Both the appendixes could be separated at the bases and were ligated individually. Our case presented a B2 type appendiceal duplication.² An inspection of the small bowel for the presence of Meckel’s diverticulum followed, which was negative. Histopathological examination of the surgical specimen confirmed the clinical findings. Both specimens showed identical features: appendiceal mucosa with extensive transmural chronic, active inflammation associated with suppurative peritonitis (Figure 3A,B). The patient had an uneventful recovery and was discharged on the sixth postoperative day.

### DISCUSSION

The first person to describe a case of double appendixes was Picoli in 1892. Its prevalence worldwide is 0.004% to 0.009% and 100 cases already reported worldwide,³ and it is found in 2 in 50,000 cases that have had surgery for appendicitis.⁴ In a search of scientific literature, most references to the double appendixes are found in case reports, indicating the rarity of this condition. In 1936, Cave proposed a classification system based on their anatomical location,⁵ and in 1963, Wallbridge revised this classification, and then modified Cave–Wallbridge classification was created.² Minor modifications were made until 1993 when Biermann suggested the following classification, which is used today:

- Type A: Single cecum with one appendix exhibiting partial duplication.
- Type B: Single cecum with two obviously separate appendixes.
  - B1: The two appendixes arise on either side of the ileocecal valve in a “bird-like” manner.
  - B2: In addition to a normal appendix arising from the cecum at the usual side, there is also a second, usually rudimentary, appendix arising from the cecum along the lines of the tenia at a varying distance from the first.
  - B3: The second appendix is located along the tenia of the hepatic flexure of the colon.
  - B4: The location of the second appendix is along the tenia of the splenic flexure of the colon.
- Type C: Double cecum, each bearing its own appendix and associated with multiple duplication anomalies of the intestinal tract as well as the urinary tract.
- Type D: Horseshoe anomaly of the appendix.³

Our case presented a B2 type of double appendixes, which is the most common type. According to a study by Nageswaran et al.⁵ there are no associated congenital abnormalities in this type of duplication; concealed duplication is confirmed only intraoperatively.

Type B2 duplication is the most common variation of anatomy and the most difficult to identify. Difficulty in

**FIGURE 1** Ultrasound findings. (A) Dilated appendix, (B) Non-Compressible, (C) Hyperechoic, (D) Intense vascularization
identification is because the appendix that arises from the convergence of the tenia is retrocecal and out of sight. Moreover, if an inflamed, anteriorly placed appendix is found, the retrocecal space is not usually explored. It is considered that approximately 37% of patients with duplication present with inflammation of both appendices at the time of operation; therefore, they may not recover postoperatively as expected. If signs of inflammation are present along the right paracolic gutter when the surgeon identifies an anteriorly placed appendix, careful examination of the cecal pole and retrocecal space should be subsequently performed.

Some extremely rare cases are described, such as “the triple appendix,”7 which cannot include the existing types. In 1986, Alvarado1 suggested a clinical diagnostic tool which considers the patient’s signs and symptoms as well as some laboratory values. It is used for stratifying the risk of appendicitis being present (Table 1). A score of 5 or 6 is compatible with the diagnosis of acute appendicitis; a score of 7 or 8 indicates probable appendicitis; and a score of 9 or 10 indicates a very probable appendicitis. The Alvarado score is considered to have high sensitivity and low specificity; therefore, it is useful in “catching” appendicitis. However, the score is less effective for stratifying the risk of appendicitis in children.8 After some years, this score was modified for patients 3–18 years old, and the Pediatric Appendicitis Score was created and implemented.

To briefly summarize the embryology of the normal appendix, during the fifth fetal week, a bud at the junction of the small and large bowel develops and undergoes rapid growth into a pouch. The proximal end of this pouch starts growing differentially to give rise to the cecum. The appendix first appears at the eighth week of gestation as an outpouching of the cecum. Then, it gradually rotates to a more medial location following

| TABLE 1  | Alvarado Score |
|----------|----------------|
| **Feature** | **Score** |
| Symptoms | |
| Migration of pain | 1 |
| Anorexia | 1 |
| Nausea | 1 |
| Signs | |
| Tenderness on the right lower quadrant | 2 |
| Rebound pain | 1 |
| Elevated temperature | 1 |
| Laboratory values | |
| Leukocytosis | 2 |
| Shift of white blood cell count to the left | 1 |
| **Total** | **10** |

FIGURE 2 Intraoperative finding of double appendixes

FIGURE 3 (A, B) Specimen A (H&E* ×4), Specimen B (H&E* ×4). Note *H&E: Hematoxylin & Eosin
the respective rotation of the gut, which results in the fixation of the cecum in the right lower quadrant. The normal embryogenesis of the appendix is well-known. However, there are no data regarding the causes of its duplication, and malrotation does not seem to be responsible for the pathophysiology of the condition. Cave has tried to explain the pathogenesis of duplication. To achieve this, he concluded with two theories: (a) the persistence of a transient embryological structure and (b) incidental appendical duplicity to a more general affection of the primitive midgut. However, even though Cave’s theories may explain some types of duplication, they are inadequate to explain all types reported.

A double appendix can be asymptomatic or cause symptoms due to obstruction or inflammation, even after an appendectomy to remove one of the two appendices. Concomitant abnormalities or duplications of the large intestine or the genitourinary system may be present in children, notably in types B1 and C, which may act as “warning” indicators due to their comparable embryological origin. The effective identification and removal of both appendices are crucial for treating this illness. Type B duplications, particularly variants in which the second appendix is located retroceally, have the highest likelihood of going unreported. Due to the increased likelihood of perforation, which can lead to widespread peritonitis, an undiagnosed second appendix might have major clinical and medicolegal consequences. A history of appendectomy in a patient with a “missed” second appendix who presents with lower abdominal pain could reasonably shift differential diagnosis to other medical conditions, such as diverticulum of the cecum, Meckel’s diverticulum, colonic adenocarcinoma, gastroenteritis, acute mesenteric adenitis, intussusception, inflammatory bowel disease, and genitourinary pathology, delaying diagnosis and appropriate laparotomy and laparoscopy have both been used successfully in the therapy of such situations. However, in circumstances where only one of the appendices is inflamed, it is critical to remove both.

It is difficult to diagnose double appendixes using routine imaging examinations. Abdominal ultrasound and computed tomography (CT) are the main methods available. However, these image examinations—mainly the CT scan—are usually not included in the routine workup of otherwise healthy patients with pain in the right iliac fossa. In cases where the patient is a child, the situation is even more challenging since a CT is avoided as the initial imaging choice due to the risk of radiation exposure. The CT has been reported to identify the double appendixes, especially in cases where both appendixes are significantly inflamed.

4 CONCLUSION

The double appendixes are rare, and its treatment is challenging. Therefore, every surgeon must be aware of the anatomical variations of the vermiform appendix, and the cecum requires routine visual inspection to ensure that there are no appendical anomalies. Double appendixes should be considered in the differential diagnosis for patients with pain in the right iliac fossa, even if the patient reports previously having an appendectomy, especially if a CT scan has not been performed. Misdiagnosis of this situation can lead to life-threatening complications for the patient and medicolegal consequences.

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CONFLICTS OF INTEREST

None.

AUTHORS CONTRIBUTIONS

This work was carried out in collaboration between all authors. Authors M. Markou and R. Sotiropoulou examined the patient, established diagnosis and participated in the surgery. Author M. Markou wrote the first draft. Author R. Sotiropoulou helped in writing. Author S. Perente participated in the surgery and reviewed the manuscript. Author I. Kougjoumtzi reviewed the manuscript. Authors S. Foutzitzi and P. Chatzipantelis established diagnosis and searched the literature. Author S. Botaitis participated in the surgery and wrote the final version of the manuscript. All authors read and approved the final manuscript.

ETHICAL APPROVAL

None.

CONSENT

Written informed consent was obtained from the patient to publish this report in accordance with the journal’s patient consent policy.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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