

Double appendix: A rare case entity

ABSTRACT

Introduction: Duplication of the vermiform appendix is a rare congenital abnormality and the majority of cases were diagnosed incidentally. **Presentation of the Case:** A forty year old male was admitted to the emergency department complaining of abdominal pain, nausea, and anorexia. The physical examination confirmed the presence of pain at Mc Burney's point. Laboratory findings showed raised WBC counts. Abdominal ultrasound revealed non compressible appendix with peri appendiceal fluid collection and probe tenderness. Intraoperatively double appendix was found .One appendix was grossly normal and other was inflamed. Both appendices were surgically removed to prevent complications in later life of the patient. **Discussion:** Although the finding of appendiceal duplication is an uncommon congenital abnormality, misdiagnosis and mismanagement of this situation may yield poor clinical outcomes and serious medical consequences. **Conclusion:** Anatomical variations should be considered as a precursor of future pathological processes.

KEY WORDS: Appendix, acute abdomen, double appendix,

INTRODUCTION

Duplication of vermiform appendix is a rare congenital anomaly originally observed and described by Picoli in 1892. Double appendix emerged as a difficult entity to diagnose. With an incidence varying from 0.004% to 0.009%^{1,2} double appendix has become a neglected and commonly missed diagnosis, often with life threatening

consequences. Double appendix are usually asymptomatic. Symptoms are usually the result of obstruction and inflammation. Most of the cases of vermiform appendix duplication in adults published in literature were incidently diagnosed during an autopsy or laparotomy for a different cause. On the other hand, double appendix in children, requires additional and more meticulous work-up as it often represents a manifestation of more complex developmental intestinal, genitourinary or vertebral abnormalities.^{1,2} The clinical presentation can vary according to the location of the appendices.³

CASE REPORT

A forty year old male patient presented to the emergency department with the chief complaint of pain abdomen, nausea and anorexia since 1 day. Pain started around umbilicus and later shifted to right lower quadrant accompanied by nausea and anorexia. On physical examination of patient, tenderness at Mc Burney's point was present. Rebound tenderness was positive at Mc Burney's point. Laboratory studies showed leucocytosis with WBC counts 13000/cumm, Hb 12gm% . Renal and liver function tests within normal limits. Ultrasound abdomen showed blind ended tubular structure with diameter of 8 mm in right iliac fossa. The tubular structure was non compressible with periappendiceal fluid collection and positive probe tenderness. Diagnosis of acute appendicitis was made and patient was shifted to operation theatre for appendicectomy. Grid iron incision was at Mc Burney's point. The presence of inflamed cecal appendix in its usual position after lysis of adhesions and cecum release was identified. After further cecal mobilization another tubular structure originating from taenia coli was identified which appeared to be appendix and grossly normal. Both the tubular structures were surgically removed. Incision closed in layers. Patient discharged on fourth post operative day in good condition. Histopathology revealed both tubular structures as appendices.



Figure 1: Intraoperative finding of double appendix.



Fig.2 : Morphology of double appendix after detached from body

DISCUSSION

The first case of double cecal appendix was reported in 1892. Since then less than 100 cases have been reported worldwide⁴. Duplication is rare and may be associated with other congenital abnormalities⁵. First classification of cecal appendix was developed in 1936 by Cave⁶ modified in 1962 by Wallbridge⁷. Since then authors made several changes to it and modified classification by Cave-Wallbridge was formed which is now most widely used. Cave-Wallbridge categorizes double cecal appendix into three types: A, B and C. Type A is characterized by the presence of two cecal appendices with a common origin in a single cecum. In type B, two appendices emerge from different cecal origins from a single cecum. This type is also subdivided into B1 and B2. In subtype B1, the two appendices emerge from a single cecum, one from each side of the ileocecal valve, symmetrically. On the other hand, in subtype B2, one of the appendices is in its usual position and the second one is located alongside the taenia coli. Finally, type C is characterized by the existence of two caecum, each with a cecal appendix. Possibility of double appendix should be kept in mind in unwell patient with clinical features suggestive of acute appendicitis, but with a macroscopically normal appendix at the time of surgery. Cecum should be carefully inspected to exclude duplicate appendix. Duplicate normal appearing appendix should be resected to avoid future diagnostic confusion⁸. Missing a duplicate appendix may have serious life threatening complications for the patient as well as medicolegal problems for the surgeon.

Conclusion

Although rare but appendix duplication should be kept in mind in patients with normal appearing appendix during surgery. Intestinal, genitourinary or bone anatomic variations should be investigated in face of duplicated appendix. Thorough examination of cecum for duplicated appendix should be carried out during surgery for acute appendicitis in case of normal appearing appendix to rule of missed appendix which can lead to serious outcomes for both patients and surgeon.

References

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