Case Report

Caecum actinomycosis with acute abdomen: A case report

Bader I. Asiri, MBBSa, Ali A. Alshehri, MD b, Abdullah S. Alqahtani, MD c, Abdullah M. Albishi, MD d, Yahia I. Assiri, MD e and Esam A. Asmiri, MBBS f

aInternal Medicine, Armed Forces Hospital Southern Region, KSA
bInfectious Diseases, Armed Forces Hospital Southern Region, KSA
cGeneral Surgery, Armed Forces Hospital Southern Region, KSA
dInternal Medicine, Gastrology and Endoscopy, Armed Forces Hospital Southern Region, KSA
eRadiology, College of Medicine in King Khalid University, KSA
fInternal Medicine, Armed Forces Hospital Southern Region, KSA

Received 19 November 2019; revised 19 January 2020; accepted 20 January 2020; Available online 19 March 2020

Abstract

Abdominal actinomycosis, one of the causes of ileocaecal disorders, is usually considered when other more common clinical conditions have been excluded. Actinomycosis is a rare infectious bacterial disorder caused by the Actinomyces species. We present the case of a 38-year male Saudi soldier who presented with pain in the right iliac fossa since 4 days prior to presentation. This stabbing pain started gradually. Based on clinical examination and abdominal ultrasound findings, an appendectomy was performed. Histological examination revealed appendicular actinomycosis with lymphoid hyperplasia, serosa congestion, and filamentous bacteria in the appendicular lumen. The patient was treated with amoxicillin. During follow-up, contrast-enhanced abdominal computed tomography (CT) and magnetic resonance imaging (MRI) revealed a 4.3 x 2.9 cm thickened caecal wall. Thereafter, the patient underwent laparoscope-assisted ileocaecal resection with ileocolic anastomosis. The histological report revealed calcified food material in the diverticulum, with chronic inflammation without actinomycosis, which may have been eradicated by the previous antibiotic treatment.

Keywords: Acute abdomen; Abdominal pain; Actinomycosis; Appendectomy; Caecum actinomycosis; Ileocaecal resection

Introduction

According to general surgical services, right iliac fossa (RIF) pain is a common condition. The differential
Diagnosis is widely varied and ranges from appendicitis to urological, gynaecological, vascular, and musculoskeletal disorders. Because clinical presentation in all these is analogous, the true cause may be uncertain, especially in case of ovarian pathologies in females of reproductive age. Therefore, diagnosing appendicitis can be challenging.2,3

Abdominal actinomycosis is one of the causes of ileo-caecal lesions. It is generally deliberated in specific clinical settings or when the more common causes have been excluded or are improbable.4 Actinomycosis is an uncommon, infectious bacterial disease caused by the Actinomyces species.5 About 70% of the infections are due to either Actinomyces israelii or Actinomyces gerencseriae.6 Infection can also be caused by other Actinomyces species, as well as by Propionibacterium propionicus, which presents similar symptoms. The condition is likely to be a polymicrobial aerobic/anaerobic infection.6

Case report

A 38-year-old male Saudi soldier from a rural area presented to our hospital with RIF pain that began four days ago. He had no relevant medical history. His symptoms started gradually as a localized stabbing pain that was aggravated by movement and decreased with rest. One day prior to presentation, the pain was associated with vomiting (no blood). An examination revealed that the patient was afebrile. The body temperature was 36.1°C, and the blood pressure was 110/78 mmHg. On palpation, the abdomen was observed to be soft and lax with no tenderness at the RIF. Abdominal ultrasound revealed free minimal fluid on the RIF, with echogenic fat planes at the assumed location of the appendix. Furthermore, a tubular structure having a maximum transverse diameter of 5.4 cm was also detected. Laboratory findings were as follows: white blood cell: 8.7, red blood cell: 4.9, haemoglobin: 15.2, platelet count: 316, total bilirubin: 22.5, direct bilirubin: 5.5, alanine aminotransferase: 27U/L, aspartate aminotransferase: 30U/L, alkaline phosphatase: 48U/L, gamma-glutamyl transferase: 16U/L, C-reactive protein: 23.8mg/l, creatinine: 80 µmol/L, and blood urea nitrogen: 4.8µmol/L. An appendectomy was performed, and subsequent histopathology revealed lymphoid hyperplasia with serosa congestion and filamentous bacteria in the lumen (Actinomycosis). The patient

Figure 1: CECT shows marked thickening of the caecum (short arrow) with a narrowed lumen, and to a lesser extent, the terminal ileum (long arrow) surrounding the fat stranding.

Figure 2: CT scan of the lower abdomen shows a dense foreign body (long arrow) impeded within the caecal wall and the fat strandings (short arrow).

Figure 3: Coronal reformatted CT shows the foreign body with surrounding hypodensity representing collections, caecal wall thickening (short arrow), and fat stranding (long heads). Based on these findings, actinomycosis infection is suggested.
underwent abdominal contrast-enhanced computed tomography (CECT) and was started on amoxicillin. CT revealed extensive wall thickening in the distal ileum along with surrounded fat (Figures 1–3). One month later, while the patient was still on amoxicillin, a follow-up abdominal CT scan revealed relative regression of the wall thickening that involved the distal ileum and caecum. However, the patient still complained of abdominal pain that recurred at least once a week. Thus, he was scheduled for contrast-enhanced abdominal magnetic resonance imaging (MRI), which revealed a caecal mass with wall thickening, measuring about 4.3 × 2.9 cm. Colonoscopy revealed a polyploid mass at the appendix (caecal mass, Figure 4). A biopsy was taken and sent for histopathology and culture. Histopathology revealed a colonic mucosal fragment with ill-defined granulomatous reaction and prominent eosinophilia with no dysplasia. Fungus culture did not yield an isolate. The patient then underwent a laparoscope-assisted ileocecal resection with ileocolic anastomosis. No postoperative complications were observed (Figure 5) and histopathology report calcified food material in diverticulosis with chronic inflammation.

Discussion

Actinomycosis is an uncommon, chronic granulomatous disease caused by filamentous, gram-positive, anaerobic bacteria. A. israelii is the main causative agent in humans. Actinomycosis has a global distribution. It mainly affects the middle-aged populations and is two to four times more common in males. Actinomycetes are the normal inhabitants of the oral cavity and the gut; however, they develop pathogenicity upon invasion of breached or necrotic tissue. As the infection progresses, granulomatous tissue, massive reactive fibrosis and necrosis, abscesses, draining sinuses, and fistulas are formed. The cervicofacial area is the most frequently infected (50%), followed by the abdominal area (20%), and the thoracic area (15%–20%). In abdominal actinomycosis, the appendix and the ileocaecal region are usually involved. The infection mostly remains localized; it then spreads contiguously, disregarding tissue planes. Lymphadenopathy is not a common finding. Hematogenous dissemination is also rare. The causative agents are the normal inhabitants of the mucous lining in the nose, throat, mouth, intestinal tract, and the female reproductive system, and are not naturally harmful. These anaerobic bacteria have the ability to grow in either the absence or in reduced concentrations of oxygen. However, any injury, trauma, or surgical procedure can cause the bacterial cells to enter deeper tissues, where they normally do not exist. Because these bacteria can grow without oxygen, they can thrive in such environments, resulting in infection.
Intestinal tract actinomycosis is a rare, chronic bacterial infection of the abdominal mucosa and/or organs of the digestive tract, mainly caused by the bacterium *A. israelii*, and sometimes by other *Actinomyces* species. Actinomycosis of the caecum is a rare, chronic bacterial infection of a part of the large intestine, caused predominantly by the bacterium *A. israelii*, and to a lesser extent by other *Actinomyces* species. The diagnosis of abdominal actinomycosis is challenging and needs surgical assessment through intervention. The cases usually have vague, nonspecific clinical complaints and the most frequent symptom is abdominal pain corresponding to the site of the infected organ. The course of the disease is indolent and is similar to that of other diseases such as appendicitis, diverticulitis, colon carcinoma, Crohn’s disease, ulcerative colitis, and tubo-ovarian abscess. In the early stages, the disease is often confused with appendicitis, carcinoma caecum, tuberculosis, or amoebiasis. The preoperative diagnosis may depend mainly on the findings of imaging modalities, which often cannot discriminate between actinomycosis and malignant process, Crohn’s disease, appendicitis, diverticulitis, or tuberculosis; however, a majority of the cases can be confirmed after surgery, based on macroscopic, microscopic, and histochemical examinations of the specimen after surgical exploration.

Evidence increasingly indicates that medical therapy alone, without surgical exploration, is the main line of treatment, irrespective of the degree or severity of the infection. Treatment of actinomycosis consists of intravenous Penicillin-G for four weeks, followed by oral penicillin for six to 12 months. Although no true surgical intervention guidelines have been established, operative treatment has been pursued in patients who present with extensive necrotic tissue or large abscesses that cannot be adequately drained.

Our patient continued to receive amoxicillin till definitive pathology. No complications arising from the surgical excision and anastomosis were observed during the follow-up. The histopathology showed no actinomycosis, which may indicate that the patient responded to the antibiotic treatment.

### Source of funding

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

### Conflict of interest

There is no conflict of interest.

### Ethical approval

The case report was approved by research ethics committee at July 15, 2019 with code No. 372.

### Authors contributions

BIA and EAA conceived, collected, organized, and designed the study, participated in writing the initial and final draft of the article. AAA participated in writing the final draft and critically revised the manuscript for intellectual content. ASA, AMA, and YIA provided the pictures and participated in writing part of the initial draft. All authors have critically reviewed and approved the final draft and are responsible for the content and similarity index of the manuscript.

### References

1. Royal College of Surgeons and Association of Surgeons of Great Britain and Ireland. Commissioning guide 2014: emergency general surgery (acute abdominal pain); 2014. London.

2. Bhangu A, Søreide K, Di Saverio S, Assarsson JH, Drake FT. Acute appendicitis: modern understanding of pathogenesis, diagnosis, and management. *Lancet* 2015 Sep 26; 386(10000): 1278–1287.

3. Andersson M, Andersson RE. The appendicitis inflammatory response score: a tool for the diagnosis of acute appendicitis that outperforms the Alvarado score. *World J Surg* 2008; 32: 1643–1649.

4. Valour F, Sénéchal A, Dupieux C, Karsenty J, Lustig S, Breton P, et al. Actinomycosis: etiology, clinical features, diagnosis, treatment, and management. *Infect Drug Resist* 2014; 7: 183–197.

5. Choi MM, Beak JH, Lee JN, Park S, Lee WS. Clinical features of abdominopelvic actinomycosis: report of twenty cases and literature review. *Yonsei Med J* 2009 Aug 31; 50(4): 555–559.

6. Baierlein SA, Wistop A, Looser C, Peters T, Riehle HM, von Flüe M, et al. Abdominal actinomycosis: a rare complication after laparoscopic gastric bypass. *Obes Surg 2007 Aug 17; (18): 1123–1126.

7. Wong VK, Turmezei TD, Weston VC. Actinomycosis. *BMJ* 2011; 343: d6099.

8. Valero R, Rodrigo E, Ruiz JC, González-Cotorruelo J, Lastra P, López-Rasines G, et al. Abcess colon diverticular disease produced for Actinomyces israelii in a renal transplant recipient. *Nefrologia: publicacion Soc Esp Nefrol* 2007; 27(4): 511–513.

9. Yegué JF, Martinez SA, Sands LR, Hellinger MD. Pelvic actinomycosis presenting as malignant large bowel obstruction: a case report and a review of the literature. *Am Surg 2000;* 66: 85.

10. Wong Randolph HL, Sihoe Alan DL, Thung KH, Wan Innes YP, Ip Margaret BY, Yim Anthony PC. "Actinomycosis: an often forgotten diagnosis". *Asian Cardiovasc Thorac Ann 2004;* 12(2): 165–167.

11. Mabeza GF, Macfarlane J. Pulmonary actinomycosis. *Eur Respir J 2003;* 21(3): 545–551. ERS Journals Ltd.

12. Acquaro P, Tagliabue F, Confalonieri G, Faccoli P, Costa M. Abdominal wall actinomycosis simulating a malignant neoplasm: case report and review of the literature. *World J Gastrointest Surg 2010;* 2: 247–250.

13. Filipou D, Psimitis I, Zizi D, Rizos S. A rare case of ascending colon actinomycosis mimicking cancer. *BMC Gastroenterol 2005 Dec;* 5(1): 1.

14. Huang CJ, Huang TJ, Hsieh JS. Pseudo-colonic carcinoma caused by abdominal actinomycosis: report of two cases. *Int J Colorectal Dis 2004;* 19: 283–286.

15. Brook I. Actinomycosis: diagnosis and management. *South Med J 2008;* 101(October 10): 1019–1023.

16. Nozawa H, Yamada Y, Muto Y, Arita S, Aisaka K. Pelvic actinomycosis presenting with a large abscess and bowel stenosis with marked response to conservative treatment: a case report. *J Med Case Rep 2007 Dec;* 1(1): 141.

17. Pusiol T, Morichetti D, Pedrazzani C, Ricci F. Abdominal-pelvic actinomycosis mimicking malignant neoplasm. *Infect Dis Obstet Gynecol 2011;* 2011: 4.
18. Aequaro P, Tagliabue F, Confalonieri G, Faccioli P, Costa M. Abdominal wall actinomycosis simulating a malignant neoplasm: case report and review of the literature. *World J Gastrointest Surg* 2010; 2: 247–250.

19. Garner JP, Macdonald M, Kumar PK. Abdominal actinomycosis. *Int J Surg* 2007; 5(6 December): 441–448 [Epub 2006 August 10].

20. Felekouras E, Menenakos C, Griniatsos J, Deladetsima I, Kalaxanis N, Nikiteas N, et al. Liver resection in cases of isolated hepatic actinomycosis: case report and review of the literature. *Scand J Infect Dis* 2004 Jul 1; 36(6–7): 535–538.

21. Bone infections. MedlinePlus. US National Library of Medicine. Retrieved 16 August 2015.

22. Islam T, Athar MN, Athar MK, Usman MHU, Misbah B. Hepatic actinomycosis with infiltration of the diaphragm and right lung: a case report. *Can Respir J Can Thorac Soc* 2005; 12(6): 336–337 [PubMed].

23. Lall T, Shehab TM, Valenstein P. Isolated hepatic actinomycosis: a case report. *J Med Case Rep* 2010 Dec; 4(1): 45.

How to cite this article: Asiri BI, Alshehri AA, Alqahtani AS, Albishi AM, Assiri YI, Asmiri EA. Caecum actinomycosis with acute abdomen: A case report. *J Taibah Univ Med Sc* 2020;15(2):148–152.