Unusual Presentations of Actinomycosis; Anterior Abdominal Wall and Appendix: Report of Three Cases

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ABSTRACT

Background: Primary actinomycosis of the anterior abdominal wall and appendix are very rare clinical entities. An accurate diagnosis is generally obtained by histological examination, and treatment often requires surgical resection.

Case Report: In this study we presented two cases of primary actinomycosis involving the anterior abdominal wall and a third one located in the appendix.

Conclusion: Actinomyces Israelii can involve all anatomic structures of the abdomen. Although preoperative diagnosis is difficult, the combination of surgery and antibiotic treatment results in complete treatment in the majority of cases.

Key Words: Actinomycosis, abdominal wall, appendix

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Introduction

Actinomyces israelii is a microaerophilic, anaerobic, gram-positive, filamentous bacterium. It is normally found in humans and can colonise the respiratory, gastrointestinal and genitourinary tracts. Actinomycosis (AMC) is a subacute to chronic bacterial infection characterised by contiguous spread, suppurative and granulomatous inflammation. It typically causes the formation of multiple abscesses and sinus tracts. Actinomycosis leads most frequently cervicofacial (50%), abdominal (20%) and thoracic (15%) involvements (1). Isolated abdominal wall disease is extremely rare, and to our knowledge, only 30 cases have been reported in the English literature to date (2). In this paper, we report two cases of actinomycosis of the anterior abdominal wall and a rare case of acute appendicitis caused by actinomycosis.

Case Reports

Case 1

A 30-year-old woman was admitted to our hospital with the complaint of a 2-week period of right upper quadrant swelling and pain. She had no history of weight loss, trauma, surgery, or intrauterine device. Upon admission, the physical exam revealed a non-movable mass in the right upper quadrant. She was febrile (38.5°C). Blood investigations revealed haemoglobin of 13.1 g/dL and white blood cell count (WBC) of 18,200/mm³. Serum amylase, blood electrolytes, liver function tests and chest radiography were normal. Computed tomography (CT) of the abdomen with oral and intravenous contrast identified an irregular density, measuring 5x8 cm arising from the internal abdominal wall. The lesion showed a faint contrast enhancement and mild inflammatory changes with no involvement of the intestines (Figure 1a).

A right subcostal laparotomy revealed an indurated mass, projecting from the posterior sheath of the rectus abdominis, without invasion of the peritoneum and abdominal viscera (Figure 1b). Surgical excision of the inflammatory mass was performed and primary closure of the wound was carried out.

The histopathological examination revealed an acute-on-chronic extended abscess of the right abdominal rectal muscle with heavy infiltration of chronic inflammatory cells and fibrous tissue with Gram-positive filamentous organisms (Actinomyces with typical sulphur granules) (Figure 1c). The patient was commenced on intravenous penicillin G (20 million IU/day) for 4 weeks postoperatively and treatment was prolonged with peroral application of amoxicillin at a dose of 2 g/day over the following 1-month period.

Case 2

The patient was a 34-year-old woman who presented to the emergency service with non-specific abdominal discomfort, a palpable mass in left lower quadrant and constipation. She described a minor weight loss over the last 2 months. Her vital signs were normal and she was afebrile. Physical examination revealed a visible left lower quadrant mass that was tender and fixed. Rectal and gynaecological examinations were normal. Laboratory results were unremarkable, apart from a WBC of 14,600/mm³. Based on the findings and the state of acute abdomen, operative intervention was planned for the patient.

A median laparotomy was performed. At surgical exploration, a large firm mass surrounded by desmoplastic reaction was found in the intestines and was densely adherent to the...
anterior abdominal wall in the left lower quadrant. Total excision of the mass was performed including the involved lateral and transverse oblique muscles as well as the subcutaneous tissue and the adjacent jejunal loop. The resulting defect of the abdominal wall was closed with a 20x20 cm prosthesis of vicryl mesh (Vicryl Mesh Knitted; © Ethicon, Inc., USA) Postoperatively, the patient recovered well without any complications. Histopathological diagnosis revealed actinomycosis with typical sulphur granules in the specimen. Medical treatment was commenced accordingly.

**Case 3**

An 18 year-old girl was referred to our clinic with a complaint of abdominal pain for 8 hours, with nausea, but no vomiting. There was right lower quadrant (RLQ) pain and tenderness with signs of peritonitis. Her medical history and blood investigations was unremarkable apart from a WBC of 15,200/mm³. Abdominal US demonstrated a target sign in the RLQ measuring 3.5 cm on cross-section. Operation was planned with the suspect of acute appendicitis. Intraoperatively, the appendix was oedematous without signs of perforation. Macroscopic examination revealed a massively swollen but intact appendiceal wall.

Thereafter, the diagnosis of AMC of the appendix was established histologically. Microscopic examination revealed typical sulphur granules and small superficial sinuses in the appendiceal lumen (Figure 2). Postoperative antibiotic treatment consisted of high doses of penicillin IV for 4 weeks. Eight months after the operation, there were no complications.

**Discussion**

Actinomycosis is a clinical infection caused by *Actinomyces israelii*. The organism is normally found in the mouth, upper respiratory, gastrointestinal, anal and genital tracts. *Actinomyces israelii* has low virulence potential and does not cause disease unless the normal mucosal barrier is broken (3). When this occurs, it leads to multiple abscess formation, fistula or mass lesions (4). It most commonly affects the head and neck (50%); however, almost any organ system can be involved (5). The abdomen can also be a site of presentation in nearly 20% of all patients. The appendix vermiformis and the ilioacaecal region are the most common localisations of AMC.

Isolated involvement of the abdominal wall is extremely rare. Similarly, there are only a few reports of clinical acute appendicitis due to AMC in the English literature (3). Laboratory tests and radiological imaging cannot significantly differentiate between AMC and other inflammatory or neoplastic processes. Preoperative diagnosis has been reported in only 10% of all cases (5). The differential diagnosis includes malignancies, amoeboma, inflammatory bowel diseases and abdominal muscle pathologies (5).
Diagnosis is revealed by histopathological demonstration of the characteristic sulphur granules containing filaments in 50% of the cases. These findings, however, are not pathognomonic, because other organisms such as Nocardia and Streptomyces may aggregate sulphur granules resembling those found in Actinomyces infections (7, 8). Microbiological isolation of the bacterium from bacterial cultures is usually inconclusive (9). Negative culture rates were reported to be as high as 76% (10).

In most cases, the diagnosis is often possible after surgical exploration and therefore the treatment becomes complex (9). The treatment of AMC is surgical drainage at the primary site of infection and, if a soft tissue tumor is suspected, a wide primary resection including the surrounding tissue. Following the confirmation of diagnosis, the systemic application of a high dose intravenous Penicillin (Penicillin G 10-20 million units daily) is the initial therapy of choice. After i.v. treatment for 4-6 weeks, oral penicillin (2-4 g/day) or amoxicillin should be administered for 6-12 months (11).

The rate of appendiceal localisation of AMC is about 65%; furthermore, the incidence seems to be increasing (12). AMC may cause acute or chronic inflammation in the appendix (12, 13). The infectious process is generally limited (12), and haematogenous or lymphatic dissemination is usually rare. A resected appendix infected by Actinomyces shows typical sinuses in microscopic examination (13); as a rule, however, the clinical presentation cannot be differentiated from suppurrative appendicitis (12, 13). Life-threatening complications such as abscesses and sinuses of the abdominal wall may occur if an actinomycotic inflammation of the appendix is left untreated (12). In the third case reported here, the diagnosis of primary actinomycosis of the appendix was revealed after pathological examination.

Abdominopelvic AMC is usually related with a history of trauma, abdominal surgery or intrauterine contraceptive devices (13). However, contrary to previous reports, no predisposing factors could be identified in our cases. In accordance with the literature, diagnosis could be made by histopathological examination of the specimen in all of our patients. Microscopic examination of the infected tissue should include a search for typical appearances of sulphur granules and, if possible, must be confirmed in microbiological cultures when a clinical suspicion of AMC arises.

All anatomic structures of the abdomen can be colonised by Actinomyces Israelli. Preoperative diagnosis is difficult because of the low index of suspicion, unusual presentations, and difficulty in culturing. In the majority of cases, the combination of surgery and antibiotic treatment results in complete treatment.

**References**