Actinomyces Odontolyticus with Klebsiella and Bacteroides Presenting as Facio-Cervical Abscess with Extension to Scalp and no Sinus Tract

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Abstract: Actinomycosis is a rare infectious bacterial disease caused by various Actinomyces species. Being very rare, Actinomycosis is easily misdiagnosed. Actinomyces are common commensals of the oral cavity which predispose to local infections. We report a case of a 67-year-old female who initially presented with swelling of the left sided face which later on proved to be a mixed infection with actinomycosis. CT imaging of head and neck was done, which revealed an abscess over the cervicofacial region. The abscess was drained and the pus was cultured in aerobic and anaerobic medium. Cultures grew actinomycosis with superadded Bacteroides and Klebsiella. The patient was managed with incision and drainage, as well as ceftriaxone. Actinomycosis is a rare infection in cervicofacial region, however, we should keep our differentials open always. Both clinical and microbiological diagnosis avoids serious errors in diagnosis.

Keywords: Actinomycosis, Facial Abscess, Mixed Abscess

Introduction

Actinomycosis is a bacterial infection. Actinomycosis can affect any part of body especially soft tissue and mainly involving in the cervicofacial region leading to abscess formation (Jameson et al., 2018). Involvement in cervicofacial region leads to abscess formation and can lead to osteomyelitis of mandible (Jameson et al., 2018; Sharkawy and Chow, 2019). Our patient had an abscess in the temporal and parotid region extending to left submandibular region. As Actinomycosis is a rare disease and since our patient had an abscess which encompassed the left temporal, parotid and submandibular regions, we report this case to highlight non-lymphatic regional spread.

Case Report

A 67 years old African American female with no previous medical conditions presented with left side facial swelling, which was progressive and painful. The patient reported that all of this started after she had pain in her left lower molar tooth.

On examination, patient had stable vital signs, including no febrile episodes. She had edema of the left-sided face with a fluctuant swelling in the left temporal region, as well as an indurated swelling at the left parotid region extending to the submandibular region. The rest of the general and systemic examination was unremarkable. Labs revealed a neutrophilic leukocytosis with normocytic, normochromic anemia, likely anemia of chronic disease secondary to the abscess. Blood glucose, BUN/CRT and liver function tests were normal. HIV screening (antigen/antibody testing) was negative. CT scan of the head and neck was obtained, which revealed a large abscess with surrounding cellulitis involving the left sided scalp and soft tissues of the neck/maxillofacial region. There was a thin communication between the scalp fluid collection and maxillofacial fluid collection anteriorly along the zygomatic arch. The cranial portion of the fluid collection measured 8.5x8.2x2.1 cm in the A-P orientation, extending from the mid parietal region to the skull base. Mild adenopathy in posterior triangle was present as shown in Fig. 1a and 1b.

Incision and drainage was performed and fluid grew Klebsiella Pneumoniae in aerobic cultures, which were sensitive to ceftriaxone. Actinomyces Odontolyticus and Bacteroides grew in the anaerobic culture media after 7 days of incubation and was sensitive to ceftriaxone. The patient was managed with flagyl for Bacteroides and ceftriaxone for Klebsiella and Actinomyces. During her treatment, her white blood cell count improved and her hemoglobin improved with good response in her reticulocyte count. Patient was scheduled to follow-up with an Infectious Disease specialist as an outpatient to complete treatment for a total of 6 months.
Discussion

Actinomycosis is a rare disease in the United States (Jameson et al., 2018; Volante et al., 2005; Smego and Foglia, 1998). Actinomyces are gram-positive, non-acid fast, anaerobic or microaerophilic filamentous branched bacteria which colonize the mouth, colon and vagina. They form normal commensals of oral cavity. Mucosal disruption can lead to infection in virtually any part of the body (Jameson et al., 2018). Actinomyces are very difficult to grow in culture, with less than 30% of cultures being positive (Volante et al., 2005). Most common Actinomyces isolated is A. Israelii, A. Propionica, A. Naeslundii, A. Viscosus and A. Odontolyticus which can lead to infections (Jameson et al., 2018).

Infection in Cervicofacial region is the most common manifestation of actinomycosis reported in literature, while involvement of central nervous system, thoracic, abdominal and pelvic actinomycosis occur less frequently (Kwartler, and Limaye, 1989; Könonen and Wade, 2015; Sharkawy and Chow, 2019). Dental procedures or having dental caries puts a patient at a high risk of cervicofacial involvement (Könonen and Wade, 2015).

In microbiology, final diagnosis, as with other microorganisms, depends upon culture. For making a final diagnosis it is important to isolate the bacteria in culture. Actinomyces generally grow in aerobic microaerophilic culture (Jameson et al., 2018). Growth of the microorganism is very difficult and less often do we see the growth (Yadav et al., 2002).

Tissue biopsy can also help in diagnosis especially as culture is difficult. Taking multiple sections from the lesion at different levels enhances chances of diagnosis (Lerner, 1988). Biopsy typically reveals acute or chronic inflammatory granulation tissue with infiltration by neutrophils, foamy macrophages, plasma cells and lymphocytes. Sulfur granules may be present (Jameson et al., 2018; Sharkawy and Chow, 2019).

Management of cervicofacial actinomycosis requires a prolonged course of antibiotics. High-dose penicillin is the treatment of choice for actinomycosis (Smego and Foglia, 1998). Surgical excision and drainage of the abscess can help in management especially if the abscess is extensive (Bennhoff, 1984).

In our patient we did not suspect actinomycosis initially because the primary abscess was over the temporal region, extending behind ear and there was no draining sinus. Diagnosis was confirmed once cultures
came back positive after seven days. The main aim of reporting this case is that one should keep the differential of a cervicofacial abscess broad and to include actinomycosis as a potential diagnosis, even if it is very rare. As the treatment for actinomycosis is for a prolonged duration, it is imperative that providers be vigilant in their diagnosis. If inadequately treated, actinomycosis can lead to serious complications.

**Conclusion**

In conclusion, although it is a rare infection, actinomycosis of the head and neck represents an important diagnosis which should not be missed. Both clinical and microbiological diagnosis avoids serious errors in diagnosis.

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**Author’s Contributions**

Tanveer Mir: Design, data collection, revised manuscript.

Anita Choudhary and Anshu Wadehra: Contributed to designing, collection of data, revised article. I gave final approval.

Pragnesh Patel: Design and data collection.

Sabah Ambreen: Design and revised manuscript.

**Ethics**

There was no animal involved in this report. We got verbal consent from the patient. We are not disclosing identity of the patient.

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