Actinomycosis of the Cheek

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INTRODUCTION

Actinomycosis is a rare chronic granulomatous infection caused by actinomycyes species; it is characterized by a slow contiguous spread and suppurrative inflammation, formation of multiple abscesses and sinus tracts with possible drainage of “sulfur granules” [1]. Actinomycyes, first described by Bollinger in 1877, are gram-positive, non-acid-fast, anaerobic or facultative anaerobic bacteria which are very difficult to cultivate [2]. This commensal bacteria usually inhabits the human oral cavity, respiratory and digestive tracts, and becomes invasive when it gains entry to the submucosal or subcutaneous tissue through cutaneous or mucosal lesions [3]. Infection is more common in rural areas than in cities. Antecedent conditions that can predispose infection are: maxillofacial trauma, tooth extraction, periodontal pockets, poor dental hygiene, immunodeficiency or conditions such as osteonecrosis after radiotherapy or after orthognathic surgery. However, there have been a few cases where the focus of infection could not be specified. The various sites of infection were described: the scalp, the forehead, the nose, the paranasal sinuses, the palate, the parotid gland, the temporal bone, the lacrimal glands, the minor salivary glands, the cheeks, the lower jaw, the tongue, the lip, the larynx and the lower pharynx [4-11]. Although uncommon, and at present estimated a rare disease in Europe [12], actinomycosis is an important clinical entity because of its difficult diagnosis due to non-specific imaging findings, symptoms that can mimic other diseases and because of long-lasting treatment. Although cervicofacial actinomycosis occurs infrequently, it should be included in the differential diagnosis when images show a soft-tissue mass with inflammatory changes and an infiltrative nature in the cervicofacial area. A CT scan and MRI are not sufficient for distinguishing actinomycosis from malignant tumoral masses [13]. Actinomycosis is now uncommon in Europe and the possibility that we may face a patient with actinomycosis is therefore underestimated [12].

We report an unusual case of actinomycosis of the cheek that occurred 6 years after buccal odontogenic abscess, intraoral incision and extraction of tooth 36.

CASE REPORT

A 56-year-old male was referred to the Department of Oral Surgery because of a painless swelling in the left cheek, which he had noticed three weeks prior to the referral. Empirical antibiotic therapy prescribed by a primary care physician did not lead to the decrease in size of the swelling. The patient who dwells in an urban area, without any pets in his household, firmly denied any contact with animals or visiting rural areas in the last 6 years, he also denied any recent odontogenic symptoms or maxillofacial trauma prior to the appearance of the swelling. However, the patient did mention having a buccal abscess 6 years earlier because of a decayed tooth 36, which was then extracted. An intraoral incision was also performed. After that painful experience the patient took good care of his remaining teeth.
and visited his primary dentist on a regular basis. The patient had no record of serious disease in his medical history and no sign of immunodeficiency. The possibility of inoculation of the bacteria with eventual biting of the buccal mucosa was excluded since the patient did not have molars in the upper and lower left jaws, which were extracted earlier except tooth 36. The patient did not use partial dentures. According to that, the only possible moment of inoculation of actinomyces was 6 years earlier, at the time when tooth 36 was extracted and incision was performed because of an odontogenic abscess in the buccal region. Clinical examination revealed a facial asymmetry, with swelling of the left cheek measuring cca 2×2 cm with a slightly erythematosus overlaying skin (Figure 1). Upon palpation the mass was round, indurated and movable. No significant cervical lymphadenopathy was noted. Intraoral examination showed partially edentulous arches with no abnormality discovered, with well maintained oral hygiene (Figure 2). Routine panoramic radiograph was performed with a normal radiological finding. The patient was referred to the Cytology Department for cytological examination. Cytological finding showed subacute inflammation and was non-specific. Thus it was decided to perform an incisional biopsy under local anesthesia. The patient gave informed consent for incision, which revealed granulation tissue with discharge of yellowish “sulfur granules”.

Histopathologic examination revealed a specimen composed of granulation tissue surrounding an abscess cavity with amorphous blue masses of actinomycete colonies in a characteristic “drusa” formation.

The diagnosis of actinomycosis was confirmed by histopathologic examination showing chronic granulomatous inflammation with the presence of multiple granules of actinomyces surrounded by polymorphonuclear neutrophils (Figures 3 and 4). According to the diagnosis oral penicillin was prescribed for four months with complete resolution.

**DISCUSSION**

Actinomyces species is divided in several subgroups. Actinomycosis in humans is mainly caused by *Actinomyces israelii*. There are at least 5 types of actinomycosis based on the site of involvement. Cervicofacial actinomycosis is most frequent and accounts for around 60% of all cases; it occurs in the oral cavity and in the region of the head and neck. Actinomyces species is commensal in the oral

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**Figure 1.** Left cheek with slightly erythematous overlaying skin

**Figure 2.** Intraoral finding, partially edentulous arches with no abnormality

**Figure 3.** Biopsy specimen composed of granulation tissue surrounding an abscess cavity with “drusa” formation of actinomycete colonies (Hematoxylin and eosin, ×40)

**Figure 4.** Biopsy specimen composed of granulation tissue surrounding an abscess cavity with “drusa” formation of actinomycete colonies (Hematoxylin and eosin, ×100)
Actinomycosis often mimics other diseases because of non-specific radiological findings and non-specific symptoms that involve indurated swelling characterized by a slow progress and with the formation of multiple sinus tracts that lack response on the empirical application of antibiotics. It is important to mention that the appearance of “sulfur granules”, although a helpful sign, is not pathognomonic for actinomycosis since they can also be present in nocardiosis [20].

Lymphatogenous spread of actinomyces is rare because of the size of bacteria; cervical lymphadenopathy eventually develops in the late stages of the disease, which can be helpful in the differential diagnosis from malignant neoplasms. Because of the above given reasons and due to a difficult cultivation of bacteria, the diagnosis of actinomycosis still remains difficult. The disease is therefore often misdiagnosed as tumorous, granulomatous disease or fungal infection [13, 21]. In obtaining the definitive diagnosis, incisional biopsy and histopathological analysis are necessary. Typical microscopical findings for actinomycosis indicate two zones; an outer zone of granulation and a central zone with multiple granules representing colonies of actinomyces [22]. Because of diagnostic difficulties, actinomycosis is also known in the literature as a great mimicker, or when referring to the cervicofacial actinomycosis as a great masquerader of head and neck disease [23, 24]. Treatment of actinomycosis consists of surgery, primarily of incision and drainage, and of antibiotic therapy. Inadequate short term antibiotic therapy can result in the relapse of infection after apparent cure. To prevent the chance of relapse of the disease, prolonged antibiotic therapy is necessary [25]. Penicillin is the antibiotic of choice, but for patients who are allergic to penicillin valid alternatives include administration of tetracycline, clindamycin, erythromycin and cephalosporin antibiotics [26].

In conclusion, we must emphasize that the prognosis of the disease is good after an early diagnosis and adequate treatment. The presented case of actinomycosis of the cheek 6 years after tooth extraction is presented as a rarity, which clinicians need to take into account in the differential diagnostics of tumors and similar lesions in the soft tissue of the oral cavity and the region of the head and neck. Because of difficult diagnosis a high dose of clinical suspicion is needed.
Актиномикоза образа

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КРАТАК САДРЖАЈ

Увод Актиномикоза је ретка хронична грануломатозна бољест коју је први пут описао Болингер 1877. године. Бољест је узрокована актиномицетама и одликује се спорим напретком, гнојном упалом, стварањем мултиплих апсцеса и фистула с могућим лучењем „сумпорних гранула“.

Приказ болесника Приказујемо необичан случај актиномикозе образа шест година након букалног одонтогеног апсцеса. Мушкарац стар 56 година упућен је у завод за оралну хирургију због безболног отока у подручју левог образа, која је започела три недеље пре упућивања у завод. Дијагноза актиномикозе потврђена је патохистопатолошким налазом. У складу са дијагнозом преписан је пеницилин током четири месеца, након чега је дошло до излечења.

Закључак Овај случај актиномикозе је ретка појава. За правилну дијагнозу потребни су пажљив преглед болесника и висок степен клиничке сумње.

Кључне речи: актиномикоза; букално; грануломатозна инфекција

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