Case report

Actinomycosis of the parotid masquerading as malignant neoplasm.

Bipin T Varghese*1, Paul Sebastian1, K Ramachandran2 and Manoj Pandey1

Address: 1Department of Surgical Oncology, Regional Cancer Centre, Trivandrum 695011, Kerala, India and 2Department of Imageology, Regional Cancer Centre, Trivandrum 695011, Kerala, India

Email: Bipin T Varghese* - geenu@eth.net; Paul Sebastian - deepakpaul@eth.net; K Ramachandran - rccvm@md2.vsnl.net.in; Manoj Pandey - manojpandey@vsnl.com

* Corresponding author

Abstract

Background: Primary actinomycosis of the parotid gland is of rare occurrence and can mimic a malignant neoplasm both clinically as well as radiologically.

Case presentation: We present here a case of primary actinomycosis of the parotid gland presenting with a parotid mass lesion with erosion of skull bones.

Conclusions: Clinical presentation of cervico-facial actinomycosis is characterized by the presence of a suppurative or indurative mass with discharging sinuses. The lesion demonstrates characteristic features on fine needle aspiration cytology and histology, however at times the findings are equivocal.

Background

Although cervico-facial actinomycosis is well described, primary actinomycosis of the parotid gland is rare. Actinomycosis was considered to be the commonest of all deep mycotic infections or mycetomas in the past [1]. However now it is well established that it is a granulomatous lesion characterized by chronic suppuration usually caused by Actinomyces israelii which is a gram positive, non-acid fast, anaerobic, commensal bacteria within the oral cavity (tonsillar crypts and tartar of teeth). Unlike most of the mycotic infections, actinomycosis is not an opportunistic infection and the portal of entry is not through inhalation. Actinomycosis usually occurs in healthy individuals when the local condition favours its growth and very often, it is a mixed bacterial growth [2]. It commonly affects the facial soft tissue although it can spread to adjacent, salivary gland, bone, and skin of face and neck. Primary actinomycosis of the parotid gland is very rare and can mimic malignancy [3].

The diagnosis of this rare condition can be reasonably made empirically even when conclusive histological or microbiological evidence is not available. A case of primary actinomycosis of the parotid gland, which was diagnosed by its clinical features, radiological findings and its response to empirical medical management, is presented.

Case presentation

A 38-year-old man presented with a 5 × 4 cm rapidly progressive, firm swelling in the left parotid region with discharging sinuses on the overlying skin. The swelling was not painful and its size did not change with meals. There was no history of fever, malaise, cough, breathlessness, haemoptysis, trauma, surgery, recent tooth extraction, or dental infection. There was no history suggestive of Hansen's disease or sexual contact. He was not diabetic or immunocompromised. The swelling was confined to the parotid region (Fig 1, 2) rest of the face including submandibular region appeared normal. A 1.5 × 1.5 cm firm
jugulodiagnostic lymph node was palpable in the neck. Oral cavity, oropharynx, and indirect laryngoscopic examinations were normal. Mouth opening was adequate and there was no facial nerve palsy. Routine investigations including blood counts, erythrocyte sedimentation rate and chest X-ray were normal. ELISA test for HIV antibody was negative. The computerized tomographic (CT) scan showed a large parotid swelling extending to the base of skull with involvement of the skull bone (Figure 3). The mass lesion was seen extending into the parapharyngeal space with thinning of the lateral wall of the maxilla on the left side (figure 4). The picture was suggestive of malignant neoplasm of the parotid gland.

Fine needle aspiration cytology (FNAC) of the parotid swelling and the neck node showed dense collections of neutrophils and macrophages with phagocytic activity, suggesting a chronic inflammatory process. Incision biopsy of the nodule with underlying parotid tissue showed salivary gland tissue with periductal lymphocytic infiltration in some of the ducts and adipose tissue, muscle bundles, sheets of lymphocytes, histiocytes, plasma
cells, neutrophils and necrosis, suggestive of chronic sia-
ladenitis. No fungal hyphae or bacteria or spores could be
demonstrated in either the FNAC or in biopsy. AFB stain-
ing was negative.

Based on the clinical, radiological and pathological find-
ings a provisional diagnosis of chronic inflammatory
pathology, probably actinomycosis was arrived at and the
patient was started empirically on oral penicillin 500 mg
6 hourly for three weeks with complete clinical response
(Fig 5, 6). The patient is disease free after and on regular
follow-up after 2 1/2 years.

Discussion
The commonest cause of a painless progressive swelling in
the parotid region is the parotid neoplasms, the other
causes being very rare (Table 1) [4-6]. A diagnosis can very
often be made by clinical and radiological features fol-
lowed by a fine needle aspiration cytology (FNAC). Actin-
omycosis of the parotid gland is rare and is often
diagnosed by the presence of bacteria or spores in either
FNAC or biopsy specimen. Microscopy may show features
of granuloma with suppuration, chronic inflammatory
cells, fibroblasts and giant cells. If suppuration is marked,
the picture is that of an acute inflammation. It is
uncommon to find colonies of microorganism in viable
tissues although they are more easily demonstrated in dis-
charge. Histological appearances are not characteristic
and not much is expected from the biopsy [1]. However it
can be diagnosed empirically, by mere exclusion, even in

Recognized routes of entry for cervico-facial actinomyco-
sis include caries tooth, tooth extraction site and tonsils. It
can also occur as a complication of trauma to respiratory
and digestive tracts including operative procedures. Infec-
tion usually starts in the subcutaneous and submucous
tissues and spreads by direct continuity [1,2].

Clinical presentation of cervico-facial actinomycosis is
characterized by the presence of a suppurative or indura-
tive mass with discharging sinuses. Pus from the discharg-
ing sinuses contains tiny yellow sulphur granules, which
are diagnostic [1]. Common initial symptoms of infection
including pain, fever, erythema, oedema, and suppura-
tion may be absent [5]. Histopathological feature are
similar to that of chronic inflammatory disease. FNAC
and exfoliative cytology may demonstrate Curshmann’s
spirals and actinomycetes filaments in fortuitous cases
[7].

Ultrasound and radiological investigations fail to differ-
entiate them from malignancy, although conventional
sialography or computerised tomography (CT) with
sialography has been suggested to present a consistent
pattern in infections [8]. A CT scan finding of soft tissue
and bony invasion, simulating malignant neoplasm,
helps in distinguishing actinomycosis from other suppu-
rative or granulomatous masses.

Drug of choice is penicillin or tetracycline. Anaerobic cul-
tures may be helpful in selecting the appropriate
antibiotic in refractory cases. Treatment is essentially medical with adjuvant surgical procedures like incision and drainage, and wound toilet [9-11]. A high index of clinical suspicion and negative tissue diagnosis of malignancy is the key to a presumptive diagnosis of this rare condition.

**Competing interest**

None declared.

**Authors contributions**

BTV carried out the literature search, prepared the draft manuscript and its subsequent revisions. PS, helped in preparing the manuscript and revised it for its scientific content. KR, did the imaging studies and contributed to the scientific content of the manuscript. MP contributed to the preparation of the manuscript and material for publication, edited the manuscript and coordinated the submission. All the authors were involved in the active patient management. All authors read and approved the final manuscript.

**Acknowledgement**

Written consent was obtained from the patient for publication of the patient’s details.

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**Pre-publication history**

The pre-publication history for this paper can be accessed here:

http://www.biomedcentral.com/1471-2407/4/7/prepub