Tuberculosis colloquativa cutis of the cheek: an extremely uncommon manifestation of primary extrapulmonary tuberculosis

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DESCRIPTION

Extrapulmonary tuberculosis (EPTB) can occur as an isolated entity or along with a pulmonary focus as in disseminated tuberculosis. Although increasingly diagnosed nowadays owing to immunodeficiency states, EPTB never fails to surprise the medical community by appearing at the rarest of the rare sites and its novel structure or appearance. The variable and myriad presentations of primary EPTB create an extremely challenging situation for the dealing physicians. However, whatever may be the presentation, it is highly gratifying to diagnose and treat this curable disease. Our case was a middle-aged, immunocompetent patient with primary tuberculous granuloma of cheek.

A 39-year-old female patient presented in the outpatient department, with right-sided facial swelling. The swelling had been gradually increasing in size for the previous 4 months with occasional bouts of pain. There were no associated nasal, ocular or dental complaints. There was no history of cough, fever, decreased appetite or weight loss. On examination, 2.5 × 2 cm firm swelling was palpable on the anterolateral wall of the right maxilla just over canine fossa, blunting the nasolabial fold. It appeared fixed to the underlying bone, but the overlying skin was mobile and smooth with no scars or sinuses. The swelling was slightly tender and did not cause any bulge in the gingivobuccal sulcus. The general physical state and the rest of the ear, nose and throat examinations were normal.

During the initial workup, fine needle aspiration cytology (FNAC) from the mass was performed; however, two FNAC reports were inconclusive and yielded only the necrotic material. Before contemplating an excision biopsy, contrast-enhanced CT of the anterior aspect of the maxillary wall was represented (figure 1). There was no evidence of maxillary bone erosion. Erythrocyte sedimentation rate was elevated, 50 mm/h (Westergren method) and the tuberculin test yielded an induration of 18 × 18 mm. Rest of the biochemical, haematological investigations were normal. Chest radiograph was within the normal limits.

The patient was taken up for excision of mass via the sublabial route under local anaesthesia. The mass was found adherent to the underlying periosteum and was removed in toto. The histopathology revealed dense inflammatory infiltrate consisting of lymphocytes, plasma cells, macrophages, multiple Langhans-type giant cells and few epithelioid cell granulomas with area of caseation necrosis and fibrosis, suggestive of tuberculosis (figure 2). Ziehl-Neelsen (ZN) stain was positive for typical acid-fast Mycobacterium tuberculosis. Postoperatively, after histopathological and microbiological confirmation of tuberculosis, the patient was started on standard multidrug antitubercular therapy. The patient has completed her therapy of 6 months duration, and is disease-free at present.

EPTB constitutes about 10% of all the cases of tuberculosis (however, this percentage may vary from region to region, in the UK ~48% of tuberculosis cases are EPTB) and cutaneous tuberculosis makes up only a small proportion (~1%) of these cases.1 2 Cutaneous tuberculosis is further classified into primary inoculation tuberculosis, tuberculosis verrucosa cutis, lupus vulgaris, tuberculosis colloquativa cutis (subcutaneous tubercular granuloma, also called as scrofuloderma), tuberculosis cutisificialis and tuberculosis cutis miliaris disseminata.3

Diagnosis of tuberculosis is difficult in such cases because of the absence of typical constitutional features. Microbiological tests such as direct microscopy and culture are usually negative in cutaneous tuberculosis because it is paucibacillary.2 Histopathological findings are characteristic but not pathognomonic and are shared by other granulomatous diseases including leprosy, sarcoidosis, leishmaniasis and subcutaneous fungal infections.1 To avoid diagnostic misinterpretation, CT scan, MRI or nuclear scans may be justified for anatomical localisation and differentiation.

Our finding of a firm soft tissue mass on the anterior aspect of the maxillary wall was representative of tuberculosis colloquativa cutis, which is a localised granulomatous tuberculous infection of

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Figure 1 Contrast-enhanced CT of the axial section of the nose and paranasal sinuses showing the soft tissue mass over the anterolateral wall of the right maxilla.
the subcutaneous tissue. The differentials considered were the inflamed lymph node, sebaceous cyst, benign growth or the granulomatous lesion. Such a case of isolated primary EPTB is extremely rare, so much so that the precise origin of the tuberculous infection can just be speculated; possibly it could be parafacial tuberculous lymphadenitis, maxillary osteomyelitis or the infected periapical tissue of maxillary molar or canine.

Though the nested PCR (nPCR) is a good tool for detecting *M. tuberculosis* DNA, in one study the correlation between nPCR results and clinical outcome was less than optimal. Therefore in granulomatous diseases of the face, it is important to evaluate not only nPCR but also the overall clinicopathological picture so as to avoid diagnostic misinterpretation. The facility for nPCR is not available at our setup, thus was not performed in the present case.

### Learning points

- Our case is a perfect example of the bizarre and atypical presentations of extrapulmonary tuberculosis (EPTB), reinstating the fact that mycobacteria can infect practically any organ of the human body.
- Diagnosis of EPTB requires high index of suspicion, even in the absence of constitutional symptoms and signs.

### Contributors

MOG is the physician and MAG is the ENT specialist who have seen and managed the case and have drafted the initial version of this manuscript. RK is the histopathologist who was involved in patient workup and management. All the authors have critically analysed the text, images and contributed significantly in shaping the final manuscript.

### Competing interests

None.

### Patient consent

Obtained.

### Provenance and peer review

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### REFERENCES


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**Figure 2** Langhans-type giant cell and epithelioid granuloma with chronic inflammatory infiltrate (H&E, ×40).