Chronic diffuse sclerosing osteomyelitis (CDSO) as stated by Jacobsson is an uncommon disease that creates diagnostic and therapeutic dilemma. A predominantly endosteal proliferative reaction of the mandible to a low-grade infection and a cyclic course with symptom-free intervals are characteristic of CDSO.[1,2] The possibility is seldom considered, simply due to rarity of the disease.[1,3]

This condition predominantly affects the body of mandible. Temporomandibular joint (TMJ) involvement with resorption, ankylosis, or osteitis of adjacent bones does not frequently occur,[4] and reports of condylar resorption in CDSO are minimal despite numerous reports of sclerosis of mandible.[1,3,5-7] Although Baltensperger demonstrated mild destruction of the condyle in 6 of the 30 assessed cases, these patients also presented with skin manifestations consistent with the diagnosis of osteomyelitis, synovitis, acne, pustulosis, hyperostosis, and osteitis (SAPHO) syndrome.[8] Similar association of condylar resorption with SAPHO syndrome has been reported in literature.[4,9,10]

Herein, we report a peculiar case of CDSO of the left hemimandible with ipsilateral condylar resorption, not associated with any dermato-skeletal conditions, as seen in the past literature. The extent of the disease is clearly illustrated by computed tomography (CT) and magnetic resonance imaging (MRI). To the best of our knowledge, condylar resorption in CDSO without underlying systemic diseases, syndromes, or multifocal skeletal defects is relatively uncommon.

**Case Report**

**Patient history**

A 37-year-old male patient reported to the Department of Oral Medicine and Radiology of this institution with a complaint of recurrent episodes of swelling and progressive trismus for the past 4 years. Each episode was followed by spontaneous remission after 2 weeks. Radiographic examination detected unilateral coarse trabeculae with ground-glass appearance of the left hemimandible and resorption of ipsilateral condyle. Computed tomography demonstrated endosteal sclerosis, cortical thickening, and condylar resorption, while magnetic resonance imaging revealed altered marrow intensity with postcontrast enhancement of the surrounding musculature. Bone histopathology was consistent with the diagnosis of osteomyelitis. Palliative antibiotic therapy and regular follow-up were recommended. Of particular interest in the present case is ipsilateral condylar resorption not associated with any dermato-skeletal conditions, which is uncommon in CDSO of the mandible.
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Post gadolinium contrast images demonstrated unilateral intramedullary enhancement, with enhancement of overlying musculature such as the left buccinator, masseter, medial and lateral pterygoid, mylohyoid, and hyoglossus muscles [Figure 5]. This suggested a diagnosis of diffuse sclerosing osteomyelitis of the left hemimandible with inflammatory changes in soft-tissue spaces and ipsilateral condylar resorption. Owing to the overall radiographic appearance, a diagnosis of diffuse sclerosing osteomyelitis was suggested.

Laboratory investigations

Erythrocyte sedimentation rate was 66 mm/h, and C-reactive protein was also elevated (11.9 mg/dl). Other blood investigations were within normal limits. Bone culture predominantly showed lymphocytes; histopathology revealed few devitalized and viable bony fragments with fibrocollagenous tissue, lymphocyte infiltration, hemorrhage, and few congested capillaries. These features were consistent with the diagnosis of chronic osteomyelitis.

Surgical treatment of such a diffuse lesion is not recommended; and hence, hyperbaric oxygen therapy was advised. However, due to financial constraints, the patient was unwilling to undergo the advised treatment. Palliative antibiotic therapy with tablet Augmentin 625 mg TDS for 5 days (Amoxicillin 500 mg + Clavulanic Acid 125 mg) as and when the episodes occur and regular follow-up was recommended.

Discussion

The term CDSO is primarily descriptive of the radiological appearance of a pathological bone reaction.[2] CDSO, also known as osteomyelitis sicca, primary chronic osteomyelitis, chronic sclerosing nonsuppurative osteomyelitis, or reactive hyperplasia of bone,[1] is a rare disease of unknown etiology, insidious in onset, and lacks an acute state.[2]

CDSO can occur at any age from 10 to 72 years[3] and is characterized by a nonsuppurative process of recurrent swelling, trismus, and dull pain.[3] It commonly affects the body of the mandible. However, Kodama et al. reported disease progression leading to the involvement of TMJ and temporal bone through extension of sclerosis from the ramus.[4] Similarly, the present case displayed extensive sclerosis along with an unusual finding of significant condylar resorption. Previous literature suggests that resorption by gradual reduction of the affected mandibular volume is a natural attempt to restore the balance between available circulatory capacity and tissue volume.[1]

The entity CDSO is reported to be a localized form of chronic recurrent multifocal osteomyelitis (CRMO) due to their identical clinical course, radiological, and histopathological features.[3,9,10] CRMO is multifocal, but mandibular involvement is uncommon, and our patient did not present with multifocal skeletal manifestations as seen in CRMO. Some authors also consider CDSO to be a mandibular localization of SAPHO syndrome, with typical features being
medullary sclerosis, subcortical erosions, and ankylosis of the condyle along with skin manifestations. \cite{4} Reports of condylar resorption associated with SAPHO syndrome have also been documented. \cite{4,9,10} The present case did not present with any dermatological manifestations. In contrast to previous reports, of particular interest in our case is the presence of condylar resorption, which is uncommon in true CDSO.

There are multiple confounding contributors toward disease progression which makes diagnostic isolation a challenge. Low virulence organisms are of etiological importance in intramedullary osseous infection causing sclerosis. \cite{1,6} However, bone cultures fail to be conclusive. \cite{1-3} In the present case, history of cyclic episodes and predomination of lymphocytes could explain the presence of a chronic low-grade infection.

CT reveals the variable degree of hyperostosis and sclerosis, which are the most striking patterns seen in CDSO. Involvement of the cortical bone results in thickening and sometimes visible loss of the cortical-medullar border, rendering cortical bone indistinguishable from cancellous bone, as seen in the present case. On MRI, edematous marrow with high-signal intensity is evident on T2-weighted images. \cite{9} CT evaluation or combination of CT with MRI is optimal, as CT provides a distinctive appearance of the sclerotic bone, while MRI defines the extent of the inflammatory process. \cite{10}

The management with long-term antibiotic therapy in early stages of the disease can have a beneficial effect by shortening...
the clinical episode. In chronic stage, however, surgical decortication combined with antimicrobial therapy might be the most effective treatment. Considering the extensive mandibular involvement in the present case, surgical treatment was not recommended as the bone in CDSO is hypovascular and healing will be protracted. Hyperbaric oxygen therapy is recommended as lowered oxygen tension and anaerobic infection are presumed. Several reports have suggested promising results with bisphosphonates as they decrease bone resorption and bone turnover. However, bisphosphonates are still far from being a standard of care in the treatment of CDSO due to several reasons, one being the risk of inducing osteonecrosis.

**Conclusion**

CDSO commonly affects the mandibular body; however, the present case demonstrated radiographic evidence of endosteal sclerosis extending up to the condylar head, leading to significant resorption. As this case presented with unspecific clinical symptoms, absence of an acute state of infection, and had an obscure etiology; it was important to review the whole course of the disease and utilize various imaging modalities to establish the correct diagnosis. Its localized occurrence with lack of association with commonly reported conditions such as CRMO and SAPHO syndrome created a diagnostic dilemma. The entity remains a diagnosis of exclusion; sufficient knowledge and awareness of the condition are imperative as it can aid in early diagnosis which perhaps will result in a less morbid intervention for resolution.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

**Financial support and sponsorship**

Nil.

**Conflicts of interest**

There are no conflicts of interest.

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