

**Intraosseous Solitary Neurofibroma in Ramus of Mandible: A Unique Clinical Case Report**

**Abstract**

Neurofibroma is a rare benign **Non**-odontogenic tumor which may occur predominantly as a feature of Von Recklinghausen's disease affecting the soft tissue. Intraorally, the intraosseous solitary variant of **neurofibroma** is a very rare entity thereby intriguing the oral physicians. We report a rare case of **solitary intraosseous neurofibroma involving the mandibular foramen on the right side of the mandible** without a family history of Von Recklinghausen's disease in a 39 year aged female **patient**. The diagnosis was made based on clinical and radiological findings and histopathological report. On **serological** investigation the patient was HIV positive. The present case is rare in regard to its location and the immunodeficiency condition of the patient.

**Introduction:**

Although the head and neck is the most common region for peripheral nerve sheath tumors, central involvement particularly in the jaw bones is quite unusual [1]. Intra oral **neurofibroma** is common in association with neurofibromatosis (Von Recklinghausen's disease), whereas solitary intraosseous neurofibroma is a rare entity since the bones do not contain myelinated nerves or nerve sheaths within their medullary spaces [2]. Among the benign nerve sheath tumors only about 45% involve the craniofacial region and in that approximately 9% occur in the oral cavity[3]and also, in the craniofacial region the most common occurrence of neurofibroma are, tongue, palate, mandibular ridge/vestibule, maxillary ridge/vestibule, buccal mucosa, lip, mandibular intrabony and gingiva[4]. The term 'solitary neurofibroma of oral cavity' was first coined by Bruce in 1954 [2]. Literature search of intraosseous neurofibroma gave a result of approximately 50 reported cases [5] including the present case, hence it can be said that occurrence of mandibular intraosseous neurofibroma is extremely rare. In this paper, we present an unusual case of intraosseous neurofibroma affecting the right side of mandible at the mandibular foramen in a 39 year old **HIV positive** female patient.

**Case report:**

A 39 year old apparently healthy female reported to us with a chief complaint of pain and swelling in the right lower one third of the face since one year. The patient experienced tingling sensation of the right half of lower lip continuously during the entire above mentioned period and was not preceded by any trauma or toothache. The swelling was initially small and gradually attained its present size. The patient gave history of pain as dull aching and intermittent in nature. Family history of the patient was non-contributory. General physical examination did not reveal any **café-au-lait** pigmentation, axial freckling or cutaneous nodules. On clinical examination, visually no gross asymmetry was found extraorally (fig.1a) or intraorally(fig.1b) .



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40 Fig. 1: Extraoral(a) and Intraoral(b) clinical photograph showing no gross asymmetry

41 The skin over the swelling was apparently normal without any scars, sinus or any drainage.  
42 Local temperature of the skin was normal. On palpation, extraorally, a bony hard well defined  
43 swelling measuring about 2x3cms was present about 2cms anterior to the pinna of the right ear  
44 and was non tender. Intraorally, swelling was palpated in the right ramus and was non tender,  
45 however, the posterior border of the swelling was inaccessible for palpation. There was no lymph  
46 node involvement in relation with the lesion.

47 The patient was advised for all routine hematological and radiological investigations along with  
48 HIV screening. The patient's routine haemogram was found to be within normal limits and the  
49 patient tested positive for HIV.

50 A panoramic radiograph, 3DCT and MRI were advised. Panoramic radiograph revealed a well  
51 circumscribed homogenously radiolucent area in the ramus of the right mandible (Fig. 2). The  
52 radiolucent area was in relation to the mandibular foramen and in continuation with the inferior  
53 alveolar nerve canal. The radiolucent area gave an appearance of a cyst within a cyst without  
54 any internal calcifications or septae.



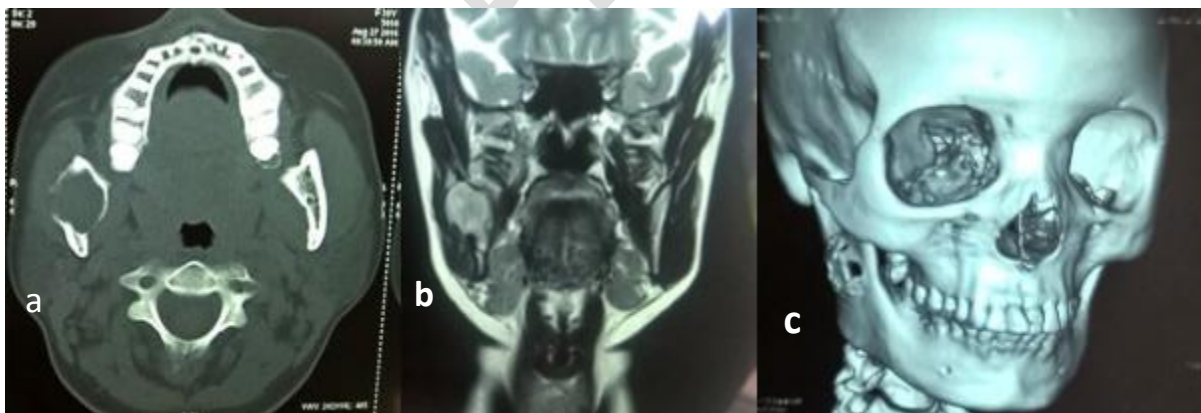
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56 Fig. 2: Orthopantomograph revealing a well circumscribed radiolucent area (white arrow)  
57 in the ramus of the right mandible

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59 3D CT scan of mandible (fig.3) revealed a well-defined expansile lytic lesion in the right  
60 mandibular ramus measuring about 2.6cmx 2.0cm x 2.6 cm with significant thinning of both  
61 medial and lateral cortices of the right mandibular ramus. Soft tissue involvement of masseter  
62 and pterygoid muscles attachment was observed.

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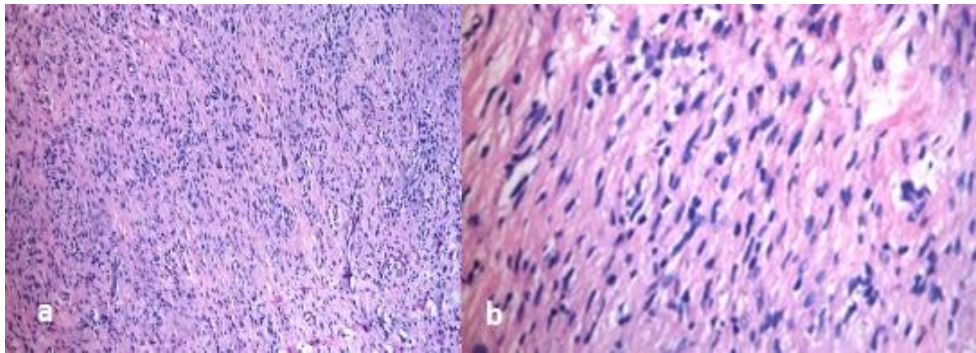
65 Fig. 3: 3DCT view (3a &3c) slice of CT scan of mandible showing a well-defined expansile lytic  
66 lesion. MRI neck (3b) showing a well-defined T2 hyperintense bone destructive lesion involving  
67 the right ramus of the mandible

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69 MRI neck (fig.3b) revealed a well-defined T2 hyperintense bone destructive lesion involving the  
70 right ramus of the mandible measuring 3.1 x 2.2 x 2.9 cms, medially the lesion abutted medial  
71 pterygoid muscle with intact fat planes and laterally the lesion abutted the inner fibers of

72 masseter muscle with loss of fat planes. Mild widening of right mental foramina with T2  
73 hyperintensity is seen along the lumen in continuous with mass lesion.

74 After an informed consent, an incisional biopsy was performed and specimen was sent for  
75 histopathological evaluation. Hematoxylin and eosin section (fig. 4) revealed a non encapsulated  
76 lesion consisting of elongated spindle cells with wavy hypochromatic nuclei dispersed in a  
77 collagenous stroma. The tumor cells are set in a fibromyxoid background interspersed with thin  
78 wirelike collagen fibres. Few areas showed whorl or fascicular arrangement of cells. Endothelial  
79 lined blood vessels along with few mast cells and diffuse chronic inflammatory infiltration was  
80 noted. A diagnosis of Solitary Neurofibroma was given.



81  
82 Fig. 4: Photomicrograph shows Hematoxylin and eosin sections with dense cellular lesion  
83 showing spindle cells with wavy nuclei and pale eosinophilic cytoplasm. (a-10X, b-40X).

84 A segment resection of the right side of the mandible was performed and the tumor was  
85 completely removed through a cervical approach under general anaesthesia(fig. 5).



86  
87 Fig. 5: Intraoperative photograph showing the surgical approach and the excised lesion

88 The postoperative course was uncomplicated and the patient was discharged. Patient was  
89 referred for counseling of HIV status and is presently under Anti Retroviral Therapy. A regular  
90 clinical follow-up is scheduled to check for any indications of any recurrence.

91 Discussion:

92 According to Polak [2] and his associates, neurofibroma of the mandible has a predilection for  
93 females (2:1) and for the posterior mandible with an average age of occurrence of 27.5 years.  
94 Neurofibroma is an uncommon benign tumor of the oral cavity often involving the trigeminal  
95 nerve which arises from the cells of the nerve sheath [6].The head and neck are commonly  
96 involved because of the rich innervations, and also superficial involvement of soft tissues is more  
97 frequent than the deeper location [7]. Most of the intraosseous neurofibromas give rise to pain  
98 and numbness of lip on the affected side due to the compression of adjacent structures by the  
99 increased tumor size. About 90% of the neurofibromas are associated with neurofibromatosis  
100 type1, and hence the presence of a solitary lesion requires physical examination and familial  
101 history to exclude Von Recklinghausen's disease [8]. The present case presented with a negative  
102 family history of neurofibromatosis, a physical examination was done and our patient did not  
103 have any clinical features suggestive of Von Recklinghausen's disease. Hence this case is unique  
104 due to the sporadic presentation of the lesion without any associated family history. Studies have  
105 shown that most intraosseous neurofibromas of the mandible are asymptomatic. So far, few cases  
106 of symptomatic intraosseous neurofibroma have been reported [8]. Larrson et al. and Apostolidis  
107 et al. reported cases of symptomatic intraosseous neurofibromas of mandible with symptoms like  
108 pain, swelling, paresthesia and bone destruction [9].Similarly, our patient presented with  
109 swelling pain and paresthesia of the lower lip on the affected side. Complete surgical excision  
110 has been the standard treatment for solitary neurofibroma and recurrence is rare [10].  
111 Interestingly, the patient was confirmed positive for HIV on investigations. HIV-1 is not a direct  
112 contributor for the occurrence of intraosseous neurofibroma. Although, literature search did not  
113 yield any case correlating the occurrence of HIV and intra osseous neurofibromas we report this  
114 case to draw attention to incidental occurrence of intraosseous solitary neurofibromas in a HIV  
115 positive patient.

116 Conclusion:

117 Radiolucent lesions in the head and neck region often pose a diagnostic challenge to the oral  
118 physians, especially those of tumors of neural origin. Although neurofibromatosis is a common  
119 occurrence, intra osseous solitary neurofibroma is a rare finding. Hence a thorough knowledge of  
120 the rarities and its varied presentation of a common disorder facilitates early diagnosis and a  
121 good prognosis, thereby providing a better quality of life to the affected individual

122 **Consent Disclaimer:**

123 As per international standard or university standard, patient's consent has been collected and preserved  
124 by the authors.

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