Notching in the Posterior Border of the Ramus of Mandible in a Patient with Neurofibromatosis Type I – A Case Report

ABSTRACT

Neurofibromatosis Type I (NFI) is a relatively common hereditary, autosomal dominant neurocutaneous condition. It is a benign peripheral nerve sheath tumour arising from Schwann cells and peripheral fibroblasts. Even though it is a neurocutaneous disorder, NF I has significant skeletal manifestations. Oral and maxillofacial skeletal structures are also often involved in the form of deepening of sigmoid notch, enlarged mandibular canal and notching of inferior border of mandible. In this paper, we are reporting a female patient of NF I who presented with a notching in the posterior border of ascending ramus of mandible.

CASE REPORT

A 38-year-old, Indo-Asian female reported to Department of Oral medicine and Radiology with chief complaint of pain in right mandibular posterior tooth. She gave no history of trauma or surgery. The patient was of short stature and had multiple skin nodules that were asymptomatic. Café a lait spots, measuring more than 7 cm, were present at multiple sites on the body. Patient’s mother and sister had similar findings. On extra-oral examination, patient revealed gross facial asymmetry due to a well defined concave defect in the right side. Also, due to this deformity, the right gonial angle was more acute as compared to the left one [Table/Fig-1]. The panoramic radiograph revealed deepening of the left coronoid notch, enlargement of bilateral mandibular canal and increased branching of the mandibular canal. A well-defined notching was present on the right posterior border of the ascending ramus [Table/Fig-2]. A clinical diagnosis of Neurofibromatosis Type I was given, based on clinical criteria of the National Institutes of Health Consensus Development Conference [1]. Since she met more than two of the minimum required criteria according to the consensus, no additional investigations were carried out.

DISCUSSION

NFI is a pleiotropic multisystem hamartomatous disorder with no sex predilection. It is caused by a mutation of the gene responsible for the production of neurofibromin, a tumour suppressor protein [1]. Individuals with NFI are at risk for skeletal abnormalities. Recent research indicated that NFI osteoprogenitor cells demonstrate increased osteoclast forming capacity, increased migration, adhesion, and in vitro bone resorption and hence, they may contribute to skeletal defects [2]. However, notching of the posterior border of the ramus of mandible, which was seen in our patient, has not been reported in literature till date. Also, recent reports suggest that most patients have tumours which are contiguous to the altered bone and hence, caution must be exercised whenever there is a bone dysplasia in a patient with NFI.
CONCLUSION
Osseous involvement in NFI is characteristic, but it differs among patients and the mandible is often affected. Knowledge on such skeletal changes is essential, since oral diagnosticians should always suspect a neoplasm which is contiguous with a bone deformity in a patient with NFI.

REFERENCES

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