Bilateral Temporal Bone Xanthomas in Type II Hypercholesterolemia

Tip 2 Hiperkolesterololemide Bilateral Temporal Kemik Ksantoması

ABSTRACT
We present a very rare case of intracranial xanthoma with bilateral temporal bone involvement. The lesion caused destruction and remodelling in mastoid air cells. On the right side, cerebellar compression was obvious. The patient was operated on the symptomatic side and the lesion was totally removed.

KEYWORDS: Bilateral, Hypercholesterolemia, Intracranial, Temporal bone, Xanthoma

ÖZ

ANAHTAR SÖZÇÜLER: Çift taraflı, Hiperkolesterololemi, İntrakranyal, Temporal kemik, Ksantoma
INTRODUCTION

Xanthomas are benign lesions which may originate from abnormal deposition of cholesterol and lipid into tissues leading to inflammatory cell infiltration (3). They affect various parts of the body; skin, tendons, flat bones, cerebral parenchym etc. (5). Intracranial xanthomas are rare entities and they may present with different symptoms depending on direct compression onto neural structures or on increased intracranial pressure (4). As far as we know, this is the first case of an intracranial xanthoma with bilateral involvement of temporal bones.

CASE REPORT

A 23-year-old male admitted with complaints of progressive headache and ataxia persisting since two years. Clinical examination showed cerebellar ataxia, positive Romberg test without any extremity weakness and hearing loss. Several xanthomas were noted on elbows, knees and over extensor surface of fingers. The patient had type II hypercholesterolemia diagnosed in childhood. The cholesterol and LDL levels were over normal ranges, 467mg/dl and 424 mg/dl, respectively. Cranial X ray and computed tomographic (CT) scans showed osteolytic lesions affecting bilateral temporal regions and destructing mastoid air cells (Figure 1A,B). On the right side, magnetic resonance imaging (MRI) scans showed heterogenous cystic lesion, containing lipid. There was no diffusion restriction on diffusion weighted images on either side (Figure 1C,D). The patient was operated on the right temporal region. After turning the skin flap, the xanthoma was visible. The lesion was lobulated, yellowish in color, filling all mastoid air cells and was destructing the temporal bone all the way down to the dura. The lesion was removed totally (Figure 2A). On histopathological examination a destructive infiltrate composed of foamy macrophages with interspersed foreign body giant cells and cholesterol clefts was demonstrated (Figure 2B).

DISCUSSION

Intracranial xanthomas are very rare entities. They generally originate from hyperlipidemic and hypercholesterolemic diseases (1, 2). They are also reported in non-hyperlipidemic states like histiocytosis, fibrous xanthomas, multicystic granulomatosis (6). Extraaxial xanthomas are generally well circumscribed lesion that may cause bony destruction and remodelling. MRI findings

Figure 1: Osteolytic nature of the xanthoma is obvious on right lateral X ray (A). The CT scan shows destruction of bilateral mastoid air cells on both temporal bones. Internal tabula of the temporal bone is also destructed on the right side (B). Preoperative T2 weighted MRI scans show bilateral lesions affecting the temporal bones. On the right side, cerebellar compression is obvious (C). On diffusion weighted image, there is hypointensity on both temporal regions with no restriction in diffusion (D).
Figure 2: Postoperative T2 weighted MR image of the patient revealed removal of the xanthoma on the right side (A). H&E photomicrograph of the xanthomatous infiltrate demonstrated the histopathological specimen characterized by foamy macrophages, foreign body giant cells and cholesterol clefts (original magnification x 100) (B).

reflect its lipid content, with heterogenous apperance being hyperintens or isointens on T1 weighted images and hypointens on T2 weighted images. There is generally neither restriction in diffusion scans nor significant contrast enhancement (4). In this location, the xanthomas should also be differentiated from cholesteatomas, cholesterol granulomas and epidermoid tumors of the temporal bone (4).

In our case, there was bilateral involvement of temporal bones. The right side was dominant and symptomatic. The symptoms relieved after removal of the lesion. General radiological apperance of the case was similar to the literature. During the surgery, this cystic component was drained. The fluid was similar to mucopurulent discharge drained from ear reported by Basavaraj et al.(3). This may correlate with degeneration of the xanthoma. In neurosurgical literature, unilateral temporal bone involvement was reported before (3, 6). However, bilateral involvement of temporal bones was not yet reported. The treatment strategy is generally total removal of the symptomatic intracranial xanthoma. We operated on this case because there was compression on the cerebellum correlating with clinical findings. The medications like resins, niacin, statins together with dietary measures may also be useful at lowering serum cholesterol levels. In selected cases, plasmapheresis may be preferred as adjunct to treatment strategy.

In conclusion, xanthomas may occupy the cranial bones bilaterally. Symptomatic ones should be removed preferentially and radically.

REFERENCES