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AJNR Am J Neuroradiol 1992, 13 (4) 1261-1264
http://www.ajnr.org/content/13/4/1261

This information is current as of August 9, 2023.
Chondromyxoid Fibroma of the Frontal Bone

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Summary: Chondromyxoid fibroma of frontal bone is a rare lesion. Plain skull films showed a round radiolucent mass with a sclerotic margin. It was dense on plain CT scan and showed no convincing contrast enhancement. MR imaging showed low signal intensity relative to gray matter on T1-weighted image (500/20), isointensity on proton-density image (2000/30), high intensity on T2-weighted image, and marked peripheral enhancement on postcontrast (gadopentetate dimeglumine) study.

Index terms: Frontal bone, fibroma; Skull, neoplasms

Fig. 1. A 41-year-old man with an indolent frontal mass. A, Plain skull films demonstrate a rounded radiolucent mass with sclerotic margin and spotty calcium deposits in the left frontal bone. B, Axial CT scans (level + 39 and window 100) show high-density lesion on plain scan (left) without convincing contrast enhancement (middle row). A bone-mode CT scan (level + 159 and window 600) shows "salt and pepper" appearance (right).
Chondromyxoid fibromas (CMF) are benign bone tumors composed of chondroid, myxoid, and fibrous elements. The majority of these tumors arise from the long bones and their overall incidence comprises less than 1% of all bone tumors (1). Those arising from the skull are rare. We have identified 13 cases reported since Jaffe and Lichtenstein established it as a histopathologic entity in 1948 (1-13). We describe a case of CMF of the frontal bone in a 41-year-old man and present the neuroradiologic findings.

Case Report

This 41-year-old man was admitted with a complaint of right-hand hypesthesia for 2 months. His past medical and family history was noncontributory. Neurologic examination revealed painless bulging of the left frontal bone that had grown slowly since the patient noted it 18 years ago. Skull films showed a round radiolucent mass with a sclerotic margin in the left frontal region that had spotty calcifications and measured 5 × 7 × 2.5 cm. (Fig. 1A) Angiography revealed dilated bilateral superior temporal and left middle meningeal arteries with a faint tumor stain. Computed tomography (CT) (Fig. 1B) showed high density in the left frontal bone on a noncontrast scan without postcontrast enhancement; bone-window views revealed a mottled appearance. Magnetic resonance (MR) (Fig. 1C), performed with a 13-cm surface-coil and a 0.5-T superconducting magnet, showed low signal intensity relative to gray matter on the T1-weighted image (500/20), isointensity on the proton-density image (2000/30), and high intensity on the T2-weighted image (2000/120), with marked

Fig. 1—Continued.
C. On sagittal MR images (upper), low signal intensity mass relative to gray matter on T1-weighted image (500/20) (left) and Gd-DTPA enhancement with central low intensity (right) were recognized. The tumor shows high intensity relative to gray matter on T2-weighted image (2000/120) (lower left), isointensity on proton image (2000/30) (lower middle), and Gd-DTPA enhancement on axial view (lower right).
Fig. 1—Continued.

D, Histologic specimen shows a pseudolobular pattern consisting of mainly myxoid (arrowheads) and fibrous (arrows) elements. Calcifications are recognized within the tumor (stars). (Hemotoxylineosin, original magnification X60.)

A left frontal craniectomy revealed a yellowish-white, soft tumor. Dissection between the tumor and the dura mater was easy and, through the dural incision, the brain parenchyma was found to be molded in the shape of the tumor. Osteoplastic cranioplasty with metacrylate plate was performed. Postoperatively, the patient has been doing well with no neurologic deficit. A decalcified tumor specimen showed a lobular pattern consisting mainly of myxoid and fibrous elements and some chondroid elements with scattered massive calcifications. (Fig. 1D) A positive reaction for Alcian blue demonstrated mucin. The histologic diagnosis was chondromyxoid fibroma.

Discussion

In 1948, Jaffe and Lichtenstein (2) described chondromyxoid fibroma, a benign bone tumor apparently derived from cartilage-forming connective tissue and composed of chondroid, myxoid, and fibrous elements (2, 14–16).

Despite its benign radiographic appearance and biologic behavior, it may be histologically misinterpreted as a malignant tumor such as chondrosarcoma because of cellular atypia. It is, therefore, one of those bone lesions in which the radiograph may be decisive in refuting the histologic impression of malignancy (17). Fechner stated that the radiographic findings are usually more diagnostic than the microscopic observations for such craniofacial fibro-osseous lesions (18).

We identified 13 cases of CMF originating from the skull in our literature review. Regarding the origin of the tumor, six arose from the frontal bone (including our case), five from the skull base, two from the parietal bone, and one from the occipital bone. Plain skull films showed radiolucent areas with irregular sclerotic margins. Calcifications were reported in five out of 10 cases that were described in detail (5, 11–13). It is of interest that all patients with skull CMF with calcification were Japanese.

CT scans were obtained in five cases. Both low- and high-density tumors with postcontrast enhancement were recognized. MR (spin-echo 1600/60) was performed only in Morikawa's case (13), where a high-intensity lesion was shown in the petrous bone, but no other documentation was available. Our case showed low intensity in relation to gray matter on the T1-weighted image. high intensity with central low intensity on the T2-weighted image, isointensity on the proton-weighted image, and positive enhancement after gadopentetate dimeglumine was given.

The treatment of choice for CMF is excision with a wide rim of normal bone. Rahimi et al (1) stated that malignant transformation is slight, unless radiation is employed, and that it need not lead to unnecessarily radical treatment.

References