CASE REPORT

PRIMARY CAVERNOUS HAEMAGIOMA OF THE SKULL: REPORT OF 2 CASES
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ABSTRACT: BACKGROUND: Cavernous haemangiomas are rare benign bone tumours and those at the level of cranial bones are even rarer. The vertebral column is most often affected, followed by the skull. Calvarial cavernous haemangioma is rare, comprising about 0.2% of all benign neoplasms of the skull. CASE REPORT: We describe here, 2 patients with calvarial cavernous haemangioma that were localized frontally and parietally. The patients were of 45 years and 50 years of age, came with the history of dizziness and headache along with slow growing mass on frontal and parietal region respectively. The mass was soft to firm in consistency with freely mobile skin above the cavernoma sites. A computed tomography scan showed an osteolytic lesion with erosion of tabula externa. Total resection of the lesions and cranioplasty were performed in both the patients. Histopathology of the surgical sample revealed primary osseous cavernous type haemangioma showing large, thin walled, dilated capillary spaces lined by flattened endothelial cells without evidence of malignancy. CONCLUSION: Skull cavernous haemangioma are rare benign tumours. It should always be considered in the differential diagnosis of malignant skull lesions. Histopathological confirmation after surgical resection of tumour is the definitive method of diagnosis.

KEYWORDS: Cavernous haemangioma, Intra-osseous tumour, Skull.

INTRODUCTION: Haemangioma are benign vascular neoplastic disorders.¹ Toynbee 1845 was the author of the first report in English, a case of haemangioma of the skull bones.² Primary intraosseous cavernous haemangioma (PICH) are benign tumours arising from intrinsic vasculature of bone.³ Primary intraosseous cavernous haemangioma are rare, benign skeletal tumours most commonly found in the spinal vertebral column.⁴ Less commonly they can involve the bones of the cranium.⁴ Haemangioma of the skull represent 0.2% of all osseous tumours and 10% of all benign tumour of the skull.¹ Typically cavernous haemangioma grow slowly before they cause symptoms of pain or a visible or palpable skull deformity.⁵

We report two rare cases of skull primary intraosseous cavernous haemangioma presented with slow growing mass on frontal region and parietal region.

Two patients one aged 45 years and other aged 50 years, presented at neurology department with history of dizziness and headache without neurological disturbances.

A 45 years old patient was having a slow growing mass on frontal region while a 50 years old man was having mass on parietal region. The masses were soft to firm in consistency with freely mobile skin above the cavernoma sites.

A computed tomography scan was performed in both the cases. It showed an osteolytic lesion with erosion of tabula externa in both the cases. Magnetic resonance imaging performed in one patient showed hypo intense lesion on T1-weighted images and hyper intense on T2-weighted image.

Craniotomy and cranioplasty had been performed and tissue sent for histopathological examination in our department.
Macroscopically both the tissues are bony fragments displayed a purple red discoloration in an area measuring 1.5 cms in diameter on outer surface. The inner table of calvaria was eroded and surrounded by bluish purple tinge.

Microscopy of both the tissues revealed few bony fragments along with large, thin walled dilated, capillary spaces lined by flattened endothelial cells without evidence of malignancy. Dilated capillary spaces are filled with blood.

Immunohistochemical studies showed CD34 positivity in the dilated vascular channels lines by endothelial cells. This finding supported the diagnosis of an intraosseous cavernous haemangioma.

DISCUSSION: Haemangiomas are benign vascular neoplastic disorders that may involve any part of the body. Histology classifies haemangioma as venous, cavernous and capillary, according to the predominant vascular network. Although cavernous haemangioma more often involve the brain parenchyma, skull bones may also be affected.

PICHs are rare, benign skeletal tumours, most commonly found in the spinal vertebral column. Cavernous haemangioma of the skull is a rare pathological diagnosis. In 1845, Toynbee first reported a case of cavernous haemangioma of the skull, and this condition was defined by Rowbotham from the histological point of view until 1924. Within the skull, calvarial haemangiomas are most common than other type of haemangiomas. In a review by Wyke, 70% of cranial PICHs were localized particularly to the parietal and frontal bones.

Heckle S, Aschoff A & Kunze S (2002) in their study done on cavernoma of the skull observed that frontal, temporal and parietal bones are the most common sites of cavernous haemangioma of skull in decreasing order. Intraosseoushaemangioma can also arise from craniomaxillofacial bones. The mandible, zygoma, maxilla & frontal bones are most commonly involved. In another review, undertaken by Barnes in 1985, the frontal bone was the most frequently involved site, followed by parietal and zygomatic process.

Our one patient had frontal while other had parietal region involvement.

PICHs commonly occur in middle aged adults in the fourth and fifth decades of life. Women are two times more commonly affected than men, although paediatric cases are also described. Our both the patients were males.

Intraosseous cavernous haemangioma is extremely benign, slow growing mostly asymptomatic tumours. As they enlarge, they present as immobile lumps on the head associated with periodic, dull throbbing headaches that occasionally develops into dizziness and severe headaches. Neurological deficits are unusual because these tumours tend to expand externally but intracranial expansion has been reported. Patients may rarely present with an associated epidural haematoma or subarachnoid haemorrhage.

Trauma seems not to be a predisposing factor in the development of these lesions. Haemangioma may be the result of faulty differentiation of primordial vessels, resulting in an abnormal capillary bed. There are several known causes for cavernous haemangioma, but some are still unknown. Studies on genes show that specific gene mutations or deletions are causes for the disease. The genes identified are KRIT1, MGC4607 and PDCD10, named CCM1, CCM2 & CCM3 respectively. The loss of function of these genes is believed to be responsible for cavernous malformations.
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Grossly, PICHs are soft rubbery, vascular tissue separated by thin bony trabeculae. Cushing described one such lesion as a purplish-black mass and mistakenly suggested as a melanotic sarcoma. Microscopically, cavernous haemangioma consist of a well-defined non-encapsulated mass, composed of large fully or partially blood filled cavernous vascular spaces, separated by scant connective tissue. Intravascular thrombosis with associated dystrophic calcification may also be seen.

Capillary haemangiomas are composed of radially directed capillary loops lined by a single layer of cuboidal epithelial cells. Capillary haemangioma may progress to cavernous.

Radiography of the skull is the most useful method to identify PICHs. In 1930 Bucy and Capp definitely described the radiographic Characteristics as a expansive, well-circumscribed area of rarefaction with a sunburst pattern of trabeculations radiating from a common center. When viewed enface or on axial views, honeycomb or soap-bubble configuration is characteristic.

CT is an excellent investigation, as it allows detailed characterization of the cortical and trabecular bone to be made. MRI investigation is important because of its potential to show soft tissue lesions. It shows isointense on T1 weighted images and hyperintense on T2 weighted images, consistent with regions of slow flowing blood. Sometime classic radiographic appearance is not evident. Consequently diagnosis is most often made during surgical resection. These tumours can be misinterpreted as lesions like multiple myeloma or osteosarcoma.

Total surgical resection is the major treatment of skull PICH. Because the imaging findings are not specific, pre-operative diagnosis is difficult, histopathology being essential.

CONCLUSION: Skull cavernous haemangioma are rare benign tumours. They do not always have typical radiologic features, so they should always be considered in the differential diagnosis of malignant skull lesions. The preferred treatment is complete tumour removal. Consequently histopathological confirmation after surgical resection is the definitive method of diagnosis.

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Fig. 1: Case 1: Haematoxylin and eosin stained sections showing intra-osseous cavernous haemangioma consisting of thin walled vascular channels lined by flattened epithelium interspersed among bony trabeculae.

Fig. 2: Case 2: Haematoxylin and Eosin stained sections showing thin walled dilated blood vascular spaces interspersed amongst bony trabeculae. Dilated capillaries are filled with blood.

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