Introduction

Osteoma is a benign bone tumor characterized by compact or cancellous bone proliferation. The aetiology of osteomas is equivocal. However, congenital anomalies, trauma and infection have been implicated.\textsuperscript{1-3} It has been reported that osteomas can occur at any age and that males and females are equally affected.\textsuperscript{4,5} However, a female predilection has also been reported.\textsuperscript{6,7} Osteomas are slow-growing lesions and patients are usually asymptomatic. However, cosmetic deformity occurs when they reach a large size and complete surgical excision may be required.\textsuperscript{4,5}

Occipital osteomas are rare.\textsuperscript{4-6} Computed tomography with 3-D volume rendering is the imaging modality of choice. Imaging with CT can help exclude involvement of the inner table of the skull and intracranial extension of tumour.\textsuperscript{4,7-10} We present a case of a slow-growing, large giant occipital osteoma in a 22-year-old female presenting with headache and cosmetic deformity of 15 years duration.

Case Report

A 22-year-old woman was referred to the Radiology Department with a complaint of occasional headache and a continuous slow-growing mass lesion on the left occipital aspect of her skull of 15 years duration. She has been aware of the slow but steady increase in the size of the lesion over the past fifteen years. She had no previous facial trauma, and her medical history was not contributory. Clinical examination was unremarkable. The lesion was hard and non-fluctuant on palpation. CT scan revealed (Fig.1-6) an inverted cup-shaped hyperdense mass lesion overlying the left occipital bone with an attenuation of 1641.9HU. It measured (15.1\times15.4\times15.6) cm in CC\times TR\times AP dimensions. The lesion had linear foci of hypodensity within it. There was no enhancement of the lesion on intravenous contrast administration. The inner table of the skull was not involved. No intracranial involvement was seen.

These clinical and radiographic findings were sufficiently supportive of the working diagnosis of giant occipital osteoma. There were no features of Gardner’s syndrome.

Due to the cosmetic deformity, the patient was
prepared for surgery. The lesion was successively excised with no post-operative sequela. The histopathologic diagnosis confirmed the clinical and radiographic diagnosis of giant occipital osteoma.

Discussion
Osteomas are rare slow-growing benign bone tumor.\textsuperscript{4,6,11} They commonly occur in the paranasal sinuses.\textsuperscript{2,13} Involvement of the temporal and occipital squama is extremely rare.\textsuperscript{2-5,14} It is frequently found in adults during the first 3 decades.
of their lives. It affects both male and female equally although other authors agree that it is mainly prevalent in women. Osteomas larger than 3cm are termed giant osteomas. They are also common in the frontoethmoidal region with above 40 cases reported in the literature. Giant occipital osteoma are rare.

Histologically osteoma can be divided into three types: mature, ivory and the mixed type. The ivory type is composed of compact bone with almost no fibrovascular interstitial tissues. It mainly occurs in the frontal sinus and has a slow growth rate. On the other hand, the mature type is composed of mature cancellous bone and mainly occurs in the maxillary sinus and ethmoid sinus. Mature type of tumor grows faster than the ivory type. Mixed osteoma is a mixture of cancellous and compact bone. The latter was reported in the index patient. Several hypothesis have been put forward for the aetiology of osteoma but were regarded as hypothesis for the early detection of Gardner’s syndrome of which osteoma is a component. Gardner syndrome is one of the polyposis syndromes. It is characterized by familial adenomyosis, multiple osteomas (especially of the mandible, skull, and long bones), epidermal cysts, fibromatoses and desmoid tumors of mesentry and anterior abdominal wall. Other abnormalities include seen in Gardner’s syndrome include supernumerary teeth, duodenal tumors /ampullary carcinoma, and papillary thyroid carcinoma.

Computed tomography is the imaging modality of choice for osteoma imaging. The main differential diagnosis of osteoma includes osteosarcoma, osteoblastic metastasis, isolated eosinophilic granuloma, ossifying fibroma, Paget's disease (sclerotic), giant cell tumor, osteoid osteoma, hemangioma, calcified meningioma, and monostotic fibrous dysplasia. Unlike malignant lesions, osteoma has a narrow zone of transition with the parent bone.

Osteomas are resected if they become symptomatic or for cosmetic reasons especially for the giant ones. The surgical target must be outlining normal cortical bone all around the lesion. Because these lesions are limited to the external cortex, finding a plane of cleavage between the osteoma and normal bone is not difficult. The prognosis of the osteoma may be considered the best in terms of cosmetic and curative aspects provided complete excision is undertaken. Malignant transformation has not been reported yet. The recurrence is also uncommon as only two cases have been reported so far.

In young patients with skull osteomas, complete workup needs to be done to rule out Gardner syndrome by screening for the concurrent presence of intestinal polyps, soft tissue tumors, and dental abnormalities.

Conclusion
We have presented a case of a giant occipital osteoma in a 22-year-old female. The lesion had grown slowly for 15 years and cause headache and cosmetic deformity. Surgical excision was done with histopathological correlation. No significant postoperative sequelae occurred.

References:
Giant osteoma of the occipital bone... 1993;22(7):485–500.