Fall while playing at home. It gradually increased in size. There was no history of headache or any neurological changes. Her neurological examination was normal. On local examination, there was a scalp swelling of about 6.0 cm × 6.0 cm × 3.0 cm at the right parasagittal region behind the coronal suture. It was noncompressible and nonpulsatile, firm in consistency with no skin discoloration. Hair distribution and temperature over the swelling was normal. There was no bruit or thrill. Cough impulse and trans-illumination test were negative. The computerized tomographic (CT) scan brain with bony windows showed a hyperdense mass involving the right parasagittal region with ground-glass appearance and expansion of involved bone [Figures 1-3]. Magnetic resonance imaging (MRI) revealed a bony defect at corresponding site with swelling at the sub-cutaneous mass with no pressure or invasion of the dura or the underlying brain parenchyma or sinus [Figure 4]. Based on history and radiological assessment, the differential diagnoses of FD and sub-periosteal calcification was made. She underwent right parasagittal craniotomy followed by cranioplasty in single sitting [Figure 5].

Her histopathology showed multiple bony trabeculae with intervening trabecular spaces showing fibroblastic proliferation and loss of periosteal rimming suggestive of FD [Figures 6 and 7]. The patient was later discharged and on 30 months follow-up there was no recurrence of the lesion with good aesthetic outcome.

## Discussion

Fibrous dysplasia is a progressive, congenital, nonheritable, benign skeletal lesion in which cellular fibrous connective tissue together with irregular woven bony trabeculae substitutes...
Figure 1: Computerized tomographic scan coronal images showing hyperdense bony mass over the right parietal bone which is approaching the midline without any parenchymal involvement

Figure 2: Computerized tomographic scan three-dimensional reconstruction showing protuberance of right parietal bone

Figure 3: Computerized tomographic scan brain plain coronal view (bone window) showing expansion of the right parietal with thinning of the cortex and ground-glass opacity

Figure 4: Magnetic resonance imaging brain contrast image showing the non-enhancing lesion of the right parietal bone approaching midline with no communication with the underlying brain parenchyma

Figure 5: Peroperative picture of resected piece of the bony mass

Figure 6: x4 microscopic view showing characteristic thin irregular curvilinear, "Chinese alphabet" spicules of woven bone separated by fibrous stroma

Figure 7: x20 microscopic view cellular fibrous tissue containing a proliferation of bland and uniform spindle cells with sparse mitotic activity with scattered thick lamellae bone without significant osteoblastic rimming focal area of normal bone.[1-3] The first reported documentation of this pathology was made by von Recklinghausen in 1891.[5] Later it was named FD by Lichtenstein in 1938.[8] It accounts for 0.8% of primary and 7% of benign bone tumors.[2]

Although the cause is unknown, there is sufficient evidence for the genetic basis of FD where mutation of the Gs α subunit of G protein-coupled receptor have resulted in proliferation of cyclic adenosine monophosphate.[7-10] In 70-80%, the disease is monostotic whereas 20-30% of cases are polyostotic.[5,7]
Monostotic condition most often involves the ribs or femur while the involvement of craniofacial bones is up to 25% of cases. This most commonly affects the maxilla and mandible followed by frontal, parietal and occipital bones.[11,12] On the other hand, polyostotic FD affects long bones, ribs, and skull, occurring unilaterally in most cases.[5] In this form there is nearly 100% involvement of the craniofacial bones with involvement of ethmoid, sphenoid, frontal, maxillary, temporal, and occipital bones in descending order of frequency.[13]

Another study of 36 patients with FD affecting skull reported that the most commonly involved area was the frontal bone (52.78%). The next was temporal bone (30.56%), followed by the sphenoid (25%), parietal bone (19.44%), and orbit (13.89%).[14]

Polyostotic FD is often seen as part of a rare condition termed McCune-Albright syndrome occurring exclusively in females, where it accompanies pigmented skin (café-au-lait spots) and endocrinopathies (sexual precocity).7,11,15 Although many patients are asymptomatic, the clinical presentation varies with the primary site of skeletal area involved, duration and extent of the disease.[13] Usually it presents as a painless slow growing mass with cosmetic disfigurement and asymmetry, headache, paresthesia or numbness, ocular proptosis, visual and hearing impairment, malocclusion, nasal obstruction, epistaxis, anosmia, epiphora, and neurological changes.[8,11,12,16]

Facial disfigurement referred to as leontiasis osseous (“lion face”) is seen with extensive involvement of the maxillary bone.[17] Majority of the patients is in their third decade at the time of diagnosis although literature has shown age range between 10 and 70 years.[17-22] Although fusion of growth plates restrict the development of any new FD lesion, however some would continue to progress despite skeletal maturation.[17] Polyostotic form of the disease affects patients at a significantly younger age. Monostotic form affects both males and females equally, but the polyostotic form is more prevalent in females.[23]

Computerized tomographic scan and MRI are essential to delineate any neurovascular and ocular involvement.[24] CT scan is the imaging modality of choice for the evaluation of FD. Characteristic findings include a “ground-glass” opacity, ballooning and expansion of the affected bone with thinning of the cortex.[25] Fries in 1957 described three different radiographic patterns of FD. These include pagetoid type (56%) which is a combination of dense and radiolucent areas of fibrosis; sclerotic type (23%), consisting of a homogenously dense mass; and cystic type (21%), with a spherical or ovoid lucency bounded by a dense rim.[26] FD is hypointense on T1-weighted image while the signal intensity on T2-weighted image varies from high to intermediate or low.

Medical management of the condition is often dealt with bisphosphonates. This reduces the risk of subsequent fracture and relieves pain.[7,26,27] Previous studies have documented the role of steroids in the treatment of an acute episode of visual impairment in patients with optic neuropathy secondary to FD.[28] For nonprogressive, asymptomatic lesions observation is generally required. Surgery is the optimum treatment modality for definitive diagnosis, disease progression or malignant degeneration, removal of compressive lesion, for cosmetic purpose and failure of nonsurgical therapy.[21,11] This varies from simple shaving, curettage or contouring of the lesion to a more aggressive resection, often accompanied by a reconstructive surgery as well to fill up the bony defect.[8,16]

It grossly appears as a firm solid grey-white mass with varied consistency and vascularity.[5,11] Histologically it is comprises of osseous and fibrous components. The osseous component is composed of thinned, irregular bone with loss of osteoblastic rimming. The intervening marrow is replaced by fibroblastic proliferation of bland spindle shaped cells.

In our case, the abnormal growth of bone started immediately after injury, however, the detailed mechanism underlying head trauma and pathogenesis of FD is still unknown, and it is possible that the relationship between these two is merely a coincidence. However, one article has reported a relationship between FD pathogenesis and increased expression of the c-fos gene.[29] Up-regulation of c-fos gene in relation to trauma has also been previously described. Therefore, there is also a possibility of induction of FD by trauma via a c-fos gene-mediated mechanism.

On follow-up the patients are clinically and radiologically assessed at regular intervals.[18] CT scan is the main stay for following such lesions after surgical excision.[1] The response to treatment can be assessed with the help of bone markers. These include total serum bone alkaline phosphatase and urine hydroxyproline, the levels of which are raised in the active phase of the disease.[7]

Spontaneous sarcomatous transformation has been reported in less than 1% of the monostotic cases mostly after radiation therapy.[30] Malignant transformation is most common to osteosarcoma,[31] although fibrosacoma, chondrosarcoma or malignant fibrous histiocytoma has also been reported.[7,9,22,28,32] Generally the outcome of FD is better except in young patients and those with polyostotic form of the disease.[7] Although rare in adults but the recurrence has been reported if the lesion is dealt surgically during their active disease phase.

**Conclusion**

To the best of our knowledge, this is the second case report of a posttraumatic cranial FD and the first case describing the lesion in the parietal bone after head injury. Most of the monostotic asymptomatic craniofacial lesions are identified incidentally. These may be surgically excised if there is a doubt in diagnosis,
cosmetic disfigurement, neurological involvement, disease progression or malignant transformation.

References