Primary Intramedullary Frontal Bone Osteoblastoma: A Case Report

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Abstract
Introduction: Osteoblastoma is a rare osteoid tissue forming tumor commonly affecting the posterior element of the spine. Intramedullary osteoblastoma of the calvarium is extremely rare. Osteoblastoma rarely occurs in the first decade of life. In this report we describe osteoblastoma of a very uncommon location in an 8-years-old male.

Case presentation: An 8-year-old male presented with a 2-year history of progressive painless swelling of the left forehead. There was no prior history of trauma to the head. Head computed tomography (CT) scan demonstrated an intramedullary oval shaped osseous lesion in the left frontal bone. Core biopsy histopathology revealed cranial osteoblastoma. Total resection of the mass was done.

Conclusion: For calvarial osteoblastoma, CT scan plays an important role in the diagnosis and differentiation from other cranial osseous tumors, yet histopathology defines the final diagnosis.

Keywords
Frontal bone, Osteoblastoma, Intramedullary, CT scan findings

Abbreviations
CT: Computed tomography

Introduction

Osteoblastoma, also known as giant osteoid osteoma, is a rare entity comprising less than 1% of all bone neoplasms [1]. Lesions widely vary in size, ranging from 2 to 10 cm with an average size of 4 cm at diagnosis [2]. It commonly affects the spine, particularly the posterior vertebral elements and the sacrum [3]. If it arises in the head, it typically affects the mandible or maxillofacial bones and, only in about 2-4% of the cases where the tumor derived from the cranial bones [4]. The temporal bone is commonly involved followed by frontal and ethmoid bones [5]. Osteoblastoma has predilection for males and mostly occur in those aged 10-30 years [6]. The lesion is often well-demarcated, with no infiltration of adjacent bone, but bone remodeling can occur depending on the size and site of the lesion. Calvarial osteoblastoma present as widened diploe with resultant thinning of the inner and/or outer tables and varying degree of calcifications [2,7]. Computed tomography (CT) scan provides information regarding bone expansion and destruction, lesion margins, host bone response, and tumor matrix [8]. Herein, we describe a rare case of primary skull bone osteoblastoma. Osteoblastoma rarely occurs in the first decade of life. In this report we describe a case of osteoblastoma in an 8-years-old male and in uncommon location that’s the frontal bone.

Case Presentation

An 8-year-old male presented with a 2-year history of a progressive yet painless left forehead swelling. Mother reported that the swelling was very small initially and grew in size with time. There was no history of trauma.
The child was relatively healthy until the swelling slowly but progressively began to emerge. The reason the mother sought medical help was not because the child felt symptoms but because the swelling was growing larger in size. Past medical and family histories were unremarkable. On clinical exam, there was a non-tender, immobile, bony hard swelling on the left forehead, 1 cm above the superior bony margin of the orbital rim that measured 3 × 2 × 0.5 cm. The overlying skin was normal and mobile over the swelling. The cough impulse and the transillumination tests were negative. There was no other swelling in the body. There was no neurologic deficit of mental status, cranial nerves, motor, sensory, deep tendon reflexes, gait and coordination and the cerebellum.

Contrasted computed tomography (CT) scan of the brain demonstrated a well-defined, expansile, oval shaped, intramedullary non-enhancing osseous lesion in the left frontal bone measured 3.9 × 2.6 × 4.0 cm. There was small intra-lesional lytic area. There was thinning of the inner plate of the cortical bone. The lesion abuts upon the orbital roof. There was no periosteal reaction. There was no intra-cranial extension. The overlying soft tissues were normal. There was no intra-conal extension. Both orbits were normal. The cerebrum, cerebellum, midbrain, pons and medulla were normal. No satellite lesions noted in the head and the demonstrated upper cervical vertebrae (Figure 1).

Core biopsy was done and the histopathology result revealed primary osteoblastoma of the cranium. Craniotomy of the outer plate of the affected frontal bone was made. Total excision of the lesion was done with great care to the inner plate (Thinning was demonstrated in the CT scan). Firm, red-brownish colored like substance was removed with ease. The inner plate, the orbital roof cortex was intact. The potential space was filled with bone cement and single plate bone flap was placed in situ. The parents and the child were satisfied with the outcome of the surgery.

Discussion

Osteoblastomas are rare entities which are benign in nature though at times aggressive osteoblastomas can present with recurrence [9]. Osteoblastomas are commonly seen in the second and third decade of life with mean age of 20 [6]. Our patient was only eight-years-old at the time of presentation.

Clinical presentation of osteoblastoma is highly dependent upon tumor location, extension, and mass ef-
fect, presentation ranges from localized pain to cranial nerve deficits and rare seizures [10]. Symptom onset is typically progressive, with an average duration of 18-24 months prior to presentation [11]. Most commonly they present as subcutaneous swelling with nocturnal pain that usually responds to salicylates [10,11]. Our patient’s frontal bone swelling was painless.

On computed tomography, calvarial osteoblastoma generally appears as a well-demarcated, mixed-lytic and sclerotic lesion with enlarged diploe, thinning outer and/or inner tables, and varying degrees of calcification [12]. At times soft tissue density can present in a portion of the lesion. It is believed that mostly it has high vascularity, and as a result can enhance [13]. Nonetheless non-enhancing lesions have also been reported [12,13], like in this case. Our patient had a widening diploe with thinning of inner table. The main portion of the lesion was an osseous lesion with central soft tissue density. Since there are differentials that have similar CT appearance like Giant cell tumor and meningioma, it should be supported by histopathology for confident diagnosis as we got in this child [14].

Osteoblastoma can grow and can be aggressive. The differentiation is mainly by histopathology though radiologically can show locally aggressive growth which often is greater than 4 cm [15]. There was no evidence of radiologic and histopathologic aggressiveness in this patient.

The treatment of choice for cranial osteoblastoma is total resection of the mass with free margins as subtotal resection can lead to greater rate of recurrence [6,16]. Chemo-radiation is reserved for unresectable lesions in cases of critical neurovascularity or recurrent lesions [16]. Our patient underwent total resection with free margins.

Conclusion

Cranial osteoblastoma is a rare entity. CT scan plays a big role in the differentiating bone tumors of the calvarium but histopathology determines the definitive diagnosis. Gross total resection is the treatment of choice; as subtotal resection has risk of recurrence.

Declaration

Ethics approval and consent to participate

The child’s parent provided an informed written consent to participate in the study.

Consent for publication

The child’s parent provided an informed written consent for this case to be published in a peer-reviewed journal.

Availability of data and materials

The information used and/or analyzed during this case report is available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Authors’ Contribution

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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References


