Unusual Large Ethmoidal Osteoma with Orbital Extension

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Introduction

Paranasal sinus osteomas are benign bone tumors. They are usually asymptomatic and found incidentally on 3% of sinus computed tomography scans [1]. They grow slowly, became symptomatic and occasionally cause complications depending on their anatomical site [1]. Osteoma with a diameter >30 mm or weighing >110 g is considered a “large” or “giant” osteoma and then surgery is required [2]. They are still more infrequent, and lead to intracranial and or orbital complications [3]. We report a rare case of a large orbito-ethmoidal osteoma, in order to analyze the epidemiological; clinical; radiological; treatment and histological aspects of this disease because the clinical characteristics and treatment remain controversial; this case report highlights the epidemiological, clinical, radiological, treatment and histological aspects of this rare disease.

Keywords: Osteoma; Benign Paranasal Pinuses Tumors; Orbit; Surgery

Report case

A 30-year-old male, presented in our ENT department; with a 2 years history of hard swelling of the medial angle of the right eye. He was in good health, and her medical history revealed recurrent rhino sinusitis. Clinical examination showed swelling in the right inner canthal area, and right painful proptosis. A significant limitation in adduction of the right globe was evident. Visual function was preserved and visual acuity was measured as 10/10 in each eye. A systemic examination was unremarkable. Nasal endoscopic examination found no nasal polyposis. Computed tomography revealed a multilobulated mass, highly calcified with osseous density, measuring 30 mm × 24 mm × 21 mm arising from right ethmoidal sinus with extension laterally in to the right orbital (Figure 1 and 2).

Abstract

Large osteomas of the fronto-ethmoidal cavities are very rare benign tumors. Headache and ocular signs are the most common symptoms. Imaging allows the diagnosis. Treatment of ethmoid sinus osteoma is surgical. One case with orbital extension is reported and discussed.

A 30 years-old male; complaining of swelling of themedial angle of the right eye. Computed tomography showed a right ethmoidal process with orbital extension. The tumor was removed through an external ethmoidectomy. Histological examination confirmed diagnosis. The patient's follow-up for the last 12 months was normal. The clinical characteristics and treatment of this disease remain controversial; this case report highlights the epidemiological, clinical, radiological, treatment and histological aspects of this rare disease.

Keywords: Osteoma; Benign Paranasal Pinuses Tumors; Orbit; Surgery

Figure 1: Coronal CT revealed a large calcified mass in the right ethmoidal sinus with orbital extension
Proctoscopy showed no abnormal findings and exclude Gardner’s syndrome. The diagnosis of a large orbito-ethmoidal osteoma was suspected. An external approach (external ethmoidectomy) was planned. The incision was Lynch-Howarth along the superomedial quadrant or region of the orbit. Dissection was continued posteriorly. Osteoma was resected en bloc using a drill and curette. The periorbital layer was preserved, and then the reconstruction wasn’t necessary. The drainage is established by widening the bone at the most dependent part in communication with the nasal cavity. The postoperative course was uncomplicated (no bleeding or cerebrospinal fluid leakage). The limitation of inward eyeball movement was improved after operation. The pathological exam confirmed diagnosis. A Computed tomography was done at six months, revealed no evidence of the osteoma (Figure 3). The patient was still free of symptoms after 12 months.

Figure 2: Axial CT revealed a large mass in the right anterior ethmoidal sinus with orbital extension

Figure 3: Postoperative axial CT scan of the patient showing total excision of the tumor (6 months after surgery)
Discussion

The orbital osteomas are particularly rare tumors representing 0.4% to 5% of all orbital tumors [4]. The most common site is the mandible, particularly the angle, followed by the sinuses. The frontal is involved in 96%, the ethmoid in 2%, and the maxillary in 2%. The sphenoid sinus is rarely affected, as are the long bones [5]. They grow slowly and when diameter became larger than 30 mm, they are considered giant tumors [2]. Osteomas affect men slightly more than women [6]. The mean age is around 50 years [7]. The etiology of osteomas is controversial. Several pathogenical hypotheses are proposed: traumatic, infective and developmental origins. Sinusitis or traumas stimulate osteoblast proliferation, leading to tumor formation. The developmental theory is explained that many osteomas arise at the junction of the ethmoid and frontal sinus, a location where membranous and cartilaginous tissues meet during embryonic life [8]. Usually, orbital osteoma is asymptomatic. In patients with giant ethmoidal sinus osteomas, the ocular signs are much more common including proptosis, diplopia and hypopsia. Also, they lead to headaches; facial deformity or fatal complications [9]. A computed tomography (CT) scan is the imaging modality of choice, because CT determines the anatomic location, the extension and the dimensions of the tumor [10]. Osteoma appears like a round or oval mass, well-circumscribed, with dense bony structure [11]. The osteomas can be observed in Gardner syndrome. This autosomal dominant disorder is characterized by intestinal polyposis, osteomas, and cutaneous and soft-tissue tumors. The risk to develop colon cancer is 100% [12]. Three histological patterns are recognised: ivory osteoma, mature osteoma, and mixed osteoma (Figure 4). Ivory osteoma also known as eburnated osteoma is a dense bone lacking haversian system. Mature osteoma or osteoma spongiosum resembles to ‘normal’ bone, including trabecular bone often with marrow. Mixed osteoma is mixture of ivory and mature histology [13]. No malignant transformation has been reported [14]. The differential diagnosis includes fibrous dysplasia, ossifying fibroma, condensing osteitis, and osteoblastoma [15]. The surgical approaches for symptomatic osteomas of the sinuses are endoscopic surgery or external approach. In asymptomatic cases, a conservative approach is recommended with regular radiological examinations [8]. Surgical treatment is recommended for:

- osteomas involving more than half of the sinus volume,
- rapidly evolving (>1 mm/year)
- Extension to the orbit or intracranial
- Location: the frontal recess and sphenoid sinus

Figure 4: histological slice showing osteoma

The surgical approach to the involved sinus is generally through a lateral rhinotomy, external fronto-ethmoidectomy, osteoplastic flap, or midfacial degloving [16,17]. The subsequent herniation of the orbital content in the ethmoid should not be considered a problem because the periorbital layer is preserved, protecting the patient from diplopia, enophthalmos, or facial deformity. Then, the reconstruction of medial orbital wall is indicated only in the case of major damage to the periorbit or in extensive intraorbital dissection. No reconstruction is required when the periorbit is spared [18]. Recently, endoscopic endonasal resection of osteomas represents a new and useful approach, offering some aesthetic advantages and lowering the morbidity noted in the classical surgical techniques. Nevertheless, this approach is not possible for voluminous orbital osteomas (like this case) [7].

Conclusion

Giant ethmoid osteomas are very rare. The ocular signs are prevalent. Treatment is surgical and the endoscopic approach offers more benefits. The rarity of orbital osteoma made our cases interesting to report

References


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