Orbital Cholesterol Granuloma: A Case Report

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Abstract

Orbital cholesterol granuloma is a rare entity with few cases described in the literature. We report the case of a 44 years old male presenting with a proptosis; an infero-medial displacement and motion limitation in upgaze of the left eye. The CT scan reveals an extraconal hypodense mass, frontal bone thinning and erosion infiltration of the zygoma. The management consists on the complete resection of the mass with curettage of the underlying bone cavity. The pathology shows cholesterol clefts; calcifications; and blood products degradation in between mononuclear cells and foamy macrophages and the absence of epithelial element thus confirming the diagnosis of cholesterol granuloma and ruling out dermoid cysts which is an important differential diagnosis.

Keywords: Proptosis; Cholesterol granuloma; Orbit curettage

Introduction

Orbital cholesterol granuloma is a rare disease with very few cases reported in the literature worldwide [1-4]. Trauma has been pointed out to be an important factor for the occurrence of this tumor [1] and cholesterol granulomas develop as an inflammatory reaction to localized hemorrhages, often occurring within the frontal bone overlying the lacrimal fossa [2]. It is usually seen in young to middle-aged males [2,3]. The aim of this work is to share a case of orbital cholesterol granuloma which should be distinguished from dermoid cyst and cholesteatoma.

Case Report

We present the case of a 44 years old male who consulted for progressive painless proptosis of left eye lasting for a year without a history of trauma. Ophthalmologic examination reveals in left eye a proptosis of 25mm; an infero-medial displacement of eye ball; limitation of eye motion in up gaze. The best corrected visual acuity; the anterior segment the fundus and the intraocular pressure were normal in both eyes. The computed tomography scan of left eye findings were: an extraconal supero-lateral hypodense mass measuring 35mm x 28mm x 25mm; frontal bone erosion with irregular border; infiltration of the zygoma bones (Figure 1).

The patient underwent a surgery by a lateral orbitotomy approach through a Reese Berke and superior lid crease incision. The tumor was completely resected and curettage of the underlying bone cavity to prevent recurrence was performed. The histopathology reveals an inflammatory granulomatous reaction with cholesterol clefts; calcifications; blood products degradation in between mononuclear cells and foamy macrophages surrounded by hyperplasia fibro-cysts and the absence of epithelial cells which are pathognomonic of orbital cholesterol granuloma (Figure 2). The outcome after 14 months follow up is favorable with absence of recurrences.
Discussion

Orbital cholesterol granuloma is a rare orbital tumor with very few reported cases [1]. It is an inflammatory reaction to foreign body. The pathogenesis is not well known but about 1/3 of reported cases point out the major role of preexisting trauma with hemorrhages occurring in the diploe of frontal bone [4]. The absence of trauma in many cases suggests the existence of others factors such as bone malformations that need to be elucidated [4,5]. It occurs mainly always in young to middle age males at the superolateral quadrant of the orbit although a case involving the superomedial quadrant of the orbit have been reported in India [5]. The computed tomography scan features are typical showing hypodense or isodense extraconal lesion in supero temporal quadrant of the orbit causing an inferomedial displacement of the eyeball, with frontal bone erosion without any calcification [5-7]. In our case we notice calcification in the mass; and infiltration in the zygoma. If for the treatment all the authors agreed that the complete excision of the mass follow by curettage of the underlying bone cavity leads to a good curative outcome [1-5], the surgical approach varies; from one to others. Some prefers the anterior orbitotomy through sub brow incision [2] others the fronto orbital craniotomy [8]. For this patient we choose the lateral orbitotomy to have a better exposure of the tumor and a well dissimilation of the incisional scar. The histopathology confirms the diagnosis by showing inflammatory cells, blood breakdown products, multinucleate giant cells; foamy macrophages and cholesterol crystals [9]. The absence of epithelial cells is a very important sign to rull out dermoid cyst and cholesteatoma [6, 9].

Conclusion

Orbital cholesterol granuloma is a rare entity. The management consists on the complete resection of the mass with curettage of the underlyng bone cavity. The pathology shows cholesterol clefts; calcifications; and blood products degradation in between mononuclear cells and foamy macrophages and the absence of epithelial element. After the surgery, the outcome is good.

References