Orbital Hemangiopericytoma Revealed by Exophthalmia: A Case Report

Essaber Hatim, Jerguigue Hounayda, Latib Rachida, Omor Youssef

1Department of Radiology, National Oncologic Institute, CHU ibn Sina Rabat, Morocco

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*Corresponding author: Essaber Hatim
Department of Radiology, National Oncologic Institute, CHU ibn Sina Rabat, Morocco

Abstract

The orbital location of hemangiopericytoma is rare, marked by its slow and unpredictable evolution with malignant potential and a high rate of recurrence and distant metastasis, hence the need for regular and prolonged monitoring, we report a case of hemangiopericytoma at man of 38 years old.

Keywords: orbital, hemangiopericytoma, exophthalmia, case, report, MRI.

INTRODUCTION

Hemangiopericytoma is a mesenchymal tumour derived from the pericyte, its orbital location is rare, it represents 0.8 to 3% of primary orbital tumours. It has an expansive character, slowly evolving, which can lead to serious ophthalmological complications.

CASE REPORT

A 38-year-old man with no notable history; who presents to the ophthalmology department for painful swelling of the left eye with ptosis evolving for a year associated with a depressive syndrome. The ophthalmological examination had objectified a limitation of ocular movements with a decrease in visual acuity of the left eye without sign of diplopia.

A cerebral CT scan was first requested and showed a spontaneously hyperdense left frontal extra-axial process with orbital extension by encompassing the optic nerve and the eyeball with grade 1 exophthalmia (Figure 1). The complement MRI, highlighted the hypervascular character by its vivid and heterogeneous enhancement after injection of Gadolinium (Figure 2).

The patient benefited from a total excision of the tumour with enucleation of the affected eye with good control after 1 year postoperatively. The anatomopathological study had confirmed the diagnosis of hemangiopericytoma.

Figure 1: Computed tomography (CT): show spontaneously hyperdense extra axial process with central zone hypodense. Extending to the left orbit

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**DISCUSSION**

Hemangiopericytoma was first described in 1942 by Stout and Murra [1] as rare vascular tumor, which originate from the proliferation of pericytes that surround the capillaries, supporting them and regulating their lumen. It generally affects the retroperitoneum and the lower limbs, its incidence in the head and neck region is less common accounts 16% and less than 5% affect the orbit [2] according to Shields’s series, only 1% of all orbital biopsies are hemangiopericytoma [3].

Most of the orbital tumors arise in one side without apparent predisposition for race or sex [2]. Orbital hemangiopericytoma almost always manifests with slowly increasing proptosis with or without associated pain and blurring or loss of vision.

The ultrasonography examine usually illustrate this tumor as an a well-defined mass with low to intermediate echogenicity [4]. On computed tomography (CT) imaging, it is seen as a well-circumscribed mass occupying either the extraconal or the intraconal space, which intensely enhances with contrast [5]. On magnetic resonance images, it is isointense in T1, hyper signal in T2 with intense enhancement.

As the clinical and radiological features are non specific, the final diagnosis of hemangiopericytoma must depend on histopathological examination of the surgical specimen.

The histopathological appearance shows tumor cells arranged in a fascicular pattern, interrupted by slit-like staghorn vessels and Immunohistochemical preparation reveals a diffuse immunoreactivity to CD34.

Its treatment is above all surgical by total excision of the tumor, the other methods are complementary such as embolization, radiotherapy, and chemotherapy to improve the local and general prognosis especially when the excision is incomplete.
CONCLUSION

The orbital location of hemangiopericytoma is rare, marked by its slow and unpredictable evolution with malignant potential and a high rate of recurrence and distant metastasis, hence the need for regular and prolonged monitoring.

REFERENCES