Primary mesenchymal chondrosarcoma of the orbit in a young female: imaging and histopathological features

Lorenzo D’Ali*, Angelica Tulisso, and Laura Mariuzzi

Department of Pathology, Ospedale Universitario Santa Maria Della Misericordia, Udine, Italy

*Correspondence address. Department of Pathology, Ospedale Universitario Santa Maria Della Misericordia, Piazzale Santa Maria della Misericordia, 15, Udine 33100, Italy. Tel: +39-0432-559401; Fax: +39-0432-559420; E-mail: lorenzo.dali88@gmail.com

Abstract

Mesenchymal chondrosarcoma (MCS) is a rare high-grade sarcoma of bone and soft tissue with highly aggressive behavior and a peak incidence in the second and third decades. We report a case of primary orbital MCS in a 30 year-old female, with radiological and clinicopathological features. Orbital MCS is an entity that should be considered in the differential diagnosis of calcified orbital lesions.

INTRODUCTION

Mesenchymal chondrosarcoma (MCS) is a rare high-grade sarcoma of bone and soft tissue firstly described by Lightenstein and Bernstein in 1959 [1].

MCS has a highly aggressive behavior and accounts for approximately 3% of all chondrosarcoma, with a peak incidence in the second and third decades [2].

MCS is composed of undifferentiated neoplastic cells associated with areas of mature cartilage and usually a hemangiopericytoma-like pattern of vascularization. Up to one-third of cases of MSC arise primarily in the soft tissue, with approximately 30 cases of orbital MSC described in the literature. Here, we report a case with primary orbital MCS in a 30 year-old patient, an entity that should be considered in the differential diagnosis of calcified orbital lesions.

CASE REPORT

A 30-year-old Caucasian female without any past medical illness presented with an endorbital mass of the right eye, without proptosis. Physical examination revealed a decreased right visual acuity. No abnormality of the left eye was detected. Computed tomography (CT) and magnetic resonance imaging (MRI) scans revealed a well-defined right intrasonic mass, in the upper-external quadrant with optical nerve displacement without involvement of it. The endorbital mass was adherent to the superior rectus muscle with a cleavage plane. CT scans demonstrated an ovoidal soft tissue mass with central calcification. T1 and T2 weighted images showed an oval tumor with iso-hyperintense signal with a calcified central component. Enhanced T2 weighted images demonstrated an omogenous tumoral enhancement except for the central calcified part (Fig. 1). These findings have led to a radiological-suggested differential diagnosis between vascular malformation, cavernous hemangioma and solitary fibrous tumor.

An excisional biopsy of the orbital mass was performed. We found a firm, whitish nodule that measured 2.8 × 1.9 × 1.7 cm. Microscopically, the neoplasm had a typical biphasic pattern, with a hypercellular small round cell component intermixed with islands of well-differentiated cartilage showing metaplastic bone ossification. There was an abrupt transition between the two different components of the neoplasm, and a hemangiopericytoma-like vascular pattern was noted...
Figure 1: (A–D) Imaging features of MCS: CT scans demonstrated an ovoidal soft tissue mass with central calcification (A), T2 (B) and T1 (C) weighted images showing an oval tumor with iso-hyperintense signal with a calcified central component. Enhanced T1 weighted image demonstrating an omogenous tumoral enhancement except for the central calcified part (D).

Figure 2: Low-power view of the lesion showing central ossification (H&E).

DISCUSSION

Clinically, the patient did not present with proptosis, a very common finding in orbital chondrosarcoma patients [3]. Similar to previous reports [4–6], we observed ossification of the orbital mass, a characteristic TC finding of orbital chondrosarcoma [4, 5].

We also observed optic nerve displacement that is a common finding in intraorbital MCS. Optic nerve tissue involvement, which has rarely been reported in huge tumors associated with intracranial extension [7], was not seen.

In our case, nuclear positivity for Sox9 was reported. This immunohistochemical finding could be useful to distinguish

(Figs 2–4). Immunohistochemically, the mesenchymal small cell component showed focal cytoplasmic positivity for CD99, while the cartilaginous component was S100 protein-positive (Figs 5 and 6). The neoplastic cells showed negativity for cytokeratin AE1/AE3, STAT6 (signal transducer and activator of transcription-6), CD31 and CD34. The findings were consistent with MCS. This case was sent for a second opinion to Prof. Angelo Paolo Dei Tos (Department of Pathology, Azienda ULSS 2Marca Trevigiana, Treviso, Italy) who confirmed the diagnosis of MCS, also reporting nuclear positivity for SOX9, a master regulator of the differentiation of mesenchymal cells into chondrocytes.
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![Image](81x607 to 261x723)

**Figure 4:** Typical hemangiopericytoma-like vascular pattern (H&E, high power view).

![Image](81x471 to 261x570)

**Figure 5:** Immunohistochemical features of MCS: cytoplasmic positivity for CD99 in the small cell component (IHC stain).

![Image](260x748)

**Figure 6:** Nuclear positivity for S100 in the cartilaginous component (IHC stain).

This entity from other small round cell tumors [8]. In addition, molecular studies have demonstrated a recurrent HEY1-NCOA2 fusion in MSC [9], which could support the diagnosis of this tumor. In conclusion, although orbital MSC is a rare entity, it should be considered in calcified lesions affecting children and young adults.

**CONFLICT OF INTEREST STATEMENT**

The authors declare that there is no conflict of interest regarding the publication of this paper.

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