Angiomyxoma of the Lower Eyelid with Orbital Extension: A Rare Entity and Review of Literature

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ABSTRACT
A 60-year-old male sought medical advice for painless progressive swelling in the left eye associated with protrusion of the eyeball and diminution of vision for the past 6 months. A computed tomography scan was done which revealed a heterogeneous lesion in the inferotemporal orbit lying close to the globe without any bony erosions or intracranial extension. Incisional biopsy revealed spindle or stellate cells in a myxoid matrix with abundant thin-walled vessels. On Immunohistochemistry, the tumour was positive for vimentin and SMA. It was negative for Desmin and S-100. A diagnosis of superficial angiomyxoma of the eyelid with orbital extension was thus made. Surgical excision under general anaesthesia was planned and the mass removed in toto. The histopathological examination of the same revealed findings similar to the incision biopsy. The patient is being followed up with no recurrences till date.

Keywords: Angiomyxoma, orbital tumor, proptosis, lid tumor, Carney’s complex

Introduction
In 1983, Steeper and Rosai described a series of distinctive soft tissue tumours of the female pelvis and perineum with the microscopic appearance of a spindle or stellate cells in a loose myxoid stroma with prominent thick-walled vessels with non-arborizing pattern and low mitotic activity. They assigned the term ‘aggressive angiomyxoma’ [1]. Allen et al., in 1988 described 30 uncommon dermal and subcutaneous angiomyxoid tumours in 28 patients. Microscopically, there were moderate to sparsely cellular angiomyxoid nodules with scattered small vessels. They coined the term ‘superficial angiomyxoma’ for such lesions.

Angiomyxomas are benign mesenchymal tumours. They rarely involve the orbit and a limited number of cases exist in literature. They can be either superficial or deep in nature. Both these varieties have been noted in orbit with one case report documenting malignant rhabdoid transformation [2]. They can be locally aggressive, which combined with their gelatinous texture precludes complete surgical removal leading to a high risk of recurrences. These recurrences usually do not cause bony erosions or tissue destruction and therefore can be managed without much morbidity. This study adhered to the tenets of the Declaration of Helsinki, and a written informed consent form was signed by the patient.

Case Report
A 60-year-old male patient presented to our outpatient department with complaints of swelling in his left lower eyelid and protrusion of the eyeball associated with diminution of vision for the past six months. (Figure 1a) On ophthalmological examination, visual acuity in right eyes was 6/6 on Snellen’s visual acuity chart and in the left eye was light perception with an inaccurate projection of rays in temporal and nasal quadrants. Both upper and lower lid fullness was noted on the left side. On Hertel’s exophthalmometry, the proptosis of the left eye was found to be 13 mm with 3 mm medial dystopia. The swelling was palpable inferiorly from the midpoint of the lower lid, laterally over the lateral canthus and superiorly till beneath the brow. The superior margin could not be felt distinctly. The mass was non-tender, firm in consistency with no palpable pulsations or thrill. It showed resistance to retropulsion with no increase on Valsalva manoeuvre. Bony orbital margins were intact. No lymph nodes were clinically palpable. Extraocular movements were found to be reduced equally in all meridians. On slit-lamp examination, inferior conjunctival prolapse was noted. Pupillary reactions were sluggish, and fundus examination using indirect ophthalmoscopy revealed disc pallor and choroidal folds at the posterior pole. Ophthalmic examination for the right eye revealed no abnormalities. Systemic examination ruled out any other lesions pertaining to Carney’s complex. A clinical differential diagnosis of lymphoma, mesenchymal tumors and orbital metastasis was made.

The patient underwent imaging, and on computed tomography, a heterogeneous lesion was noted in the inferotemporal orbit lying in close relation to the globe. (Figure 1c, d). An incisional biopsy was then planned under local infiltrative anaesthesia.
On histopathological examination of the biopsy, the lesion displayed scattered spindle-to stellate-shaped cells with ill-defined cytoplasm in the loose myxoid stroma with inflammatory cell infiltration. (Figure 2b) The cells had oval, hyperchromatic nuclei with indistinct nucleoli. (Figure 2c) There were no mitotic figures or atypical cells. The stroma was rich in collagen fibres and pale amphophilic to eosinophilic. The tumour was poorly circumscribed with infiltrative margins and displayed abundant thin-walled capillaries and low cellularity. Based on this finding, differentials of superficial anxiomyxoma, myxoid fibrous histiocytoma and dermatofibrosarcoma protuberans.

were made. On further immunohistochemical analysis, the tumour was positive for vimentin and smooth muscle actin and negative for Desmin, indicating myofibroblastic differentiation. (Figure 2d&e) S-100 staining was also noted to be negative thus ruling out Schwann cell or melanocytic origin. Ki-67 index was less than 1%, indicating low proliferative activity in the incised mass.

The mass was then removed in toto via lateral orbitotomy under general anaesthesia performed by a modified Burke-Kronlein incision. Grossly the tumour was well-circumscribed myxoid mass measuring 25 x 22 mm. (Figure 2a) The histopathological examination of the removed mass corroborated the findings of the incisional biopsy performed preoperatively.

Following the surgery, the proptosis disappeared. We have a six month follow up of the patient, and there have been no signs of recurrence to date. (Figure 1b)

**Discussion**

Anxiomyxomas of the orbit are a rare presentation. A literature search using the MESH terms ‘orbital angiomyxoma’, ‘angiomyxoma’, ‘orbital tumors’ was done on PubMed and google scholar, and we found a total of 10 cases of orbital angiomyxomas reported in literature till May 2020. (Table 1)

Though comprising a very rare group of tumours involving the orbit and periorbital cutaneous tissues, angiomyxoma, both superficial and deep/aggressive, need to be considered as differential diagnoses because these tumours are locally aggressive and incomplete removal leads to a high rate of recurrences. These lesions also had a tendency to recur, which ranged from 23% in tumours without epithelial

<table>
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<tr>
<th>Author/ year</th>
<th>Location</th>
<th>Patient Any comorbidities</th>
<th>Final diagnosis</th>
<th>Clinical features</th>
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<th>Treatment</th>
<th>IHC</th>
<th>Follow up, recurrence</th>
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</thead>
<tbody>
<tr>
<td>Hidayat et al. 2007*</td>
<td>Case 1 Left orbit, cavernous sinus, base of cranial fossa</td>
<td>7/M</td>
<td>Angiomyxoma</td>
<td>Proptosis, loss of inferior VF, decreased visual acuity</td>
<td>MRI showed a tumor that filled and expanded the left cavernous sinus, having extended from the left orbital apex, where it involved the superior and inferior rectus muscles and compressed the optic nerve.</td>
<td>Transcranial orbitotomy for orbital tumor excision, dissection of tumor from cranial nerves, methotrexate</td>
<td>Vimentin + CD34 + Factor XIIIA + SMA +/- CD68 + S-100 - Desmin -</td>
<td>5 months 1 recurrence</td>
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<td>Case 2</td>
<td>Right orbit, ethmoid and maxillary sinuses, cavernous sinus</td>
<td>4/F</td>
<td>Angiomyxoma</td>
<td>Proptosis, ptosis, afferent pupillary defect, decreased visual acuity</td>
<td>CT revealed a right posterior, subperiosteal, superomedial, homogenous mass with erosion of the medial orbital wall as well as a right maxillary sinus retention cyst. Magnetic resonance imaging showed a 30x21x15-mm mass extending to the orbital apex and displacing the optic nerve. The tumor was isodense on T1- and bright on T2- weighted images; it enhanced with contrast.</td>
<td>Frontal craniotomy, orbitotomy, ethmoid and maxillary sinuses for removal of tumor, orbital exenteration</td>
<td>Vimentin +++ CD34 +++ factor XIIIa+ SMA +/- CD68 + S-100 -, Desmin -, EMA -, GFAP -</td>
<td>8 years 3 recurrences</td>
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<td>Hidayat et al. 2007</td>
<td>Case 3</td>
<td>Right orbit</td>
<td>48/M</td>
<td>Angiomyxoma</td>
<td>Painless palpable mass</td>
<td>Not Available</td>
<td>Simple local excision</td>
<td>Vimentin ++ CD34 ++ Factor XIIIa+ SMA – Desmin -</td>
</tr>
<tr>
<td>Bajaj et al. 2011</td>
<td>Case 1</td>
<td>Left superomedial orbit</td>
<td>40/F</td>
<td>Angiomyxoma</td>
<td>Painless palpable mass, 6/36 visual acuity</td>
<td>USG suggestive of heterogeneous mass with low to medium amplitude spikes</td>
<td>Complete surgical removal by anterior orbitotomy</td>
<td>Diffuse reactivity for vimentin and focal positivity for CD34. S-100 – Cytokeratin – Desmin -</td>
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<td></td>
<td>Case 2</td>
<td>Right superomedial orbit</td>
<td>32/M</td>
<td>Angiomyxoma</td>
<td>Proptosis, occasional pain, 6/6 both eyes</td>
<td>CT scan: heterogeneous enhancing mass, with irregular borders, infiltrating surrounding tissue and LPS, SR muscles MRI: tumor iso intense on T1- and bright on T2-weighted images; enhanced with contrast</td>
<td>Debubling surgery, recurrence at 6 months, large solid mass removed via superomedial orbitotomy</td>
<td>Diffuse reactivity for vimentin and focal positivity for CD34. S-100 – Cytokeratin – Desmin -</td>
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<td>Case 3</td>
<td>Right superomedial orbit</td>
<td>28/M</td>
<td>Aggressive angiomyxoma</td>
<td>Proptosis, occasional pain, 6/6 both eyes</td>
<td>CT scan: heterogeneous enhancing mass, with irregular borders infiltrating MR, LPS, SR muscles</td>
<td>Piecemeal removal via anterior orbitotomy</td>
<td>Diffuse reactivity for vimentin and focal positivity for CD34. S-100 – Cytokeratin – Desmin -</td>
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<td>Case 4</td>
<td>Left superior orbit</td>
<td>40/M</td>
<td>Aggressive angiomyxoma</td>
<td>Proptosis, painless palpable mass, decreased visual acuity of 6/24, RAPD present</td>
<td>CT scan: heterogeneous enhancing mass, with irregular borders</td>
<td>Mass removal via antero-lateral orbitotomy</td>
<td>Diffuse reactivity for vimentin and focal positivity for CD34. S-100 – Cytokeratin – Desmin -</td>
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<td>Mishulin et al. 2012</td>
<td>Right glabellar and infra-medial brow</td>
<td>62/M</td>
<td>Aggressive Glabellar Angiomyxoma with Orbital Extension</td>
<td>Painless mass</td>
<td>CT Scan: a 2.5 x 1.8 cm glabellar cystic mass with extension to the right exacanal anterior orbit. The underlying cortical bone was intact, lacking erosion or invasion.</td>
<td>Mass excision via glabellar incision</td>
<td>NA</td>
<td>NA</td>
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<td>Jakobiec et al. 2015</td>
<td>Left medial orbit</td>
<td>47/F</td>
<td>Aggressive orbital angiomyxoma with malignant rhabdoid transformation</td>
<td>1981: painless left upper and lower eyelid swelling 2000: VA 20/20 OD, 20/30 PS, 6 mm of left-sided proptosis, diplopia on right gaze, and a palpable mass below the left medial canthus 2012: vision no light perception in left eye, frozen globe, proptosis &gt;35mm, Severe lagophthalmos with bulbous interpalpebral chemosis</td>
<td>CT scan 2000: a multiloculated mass with internal septa in the left medial orbit. The medial wall of the orbit was bowed toward the ethmoid sinus. CT scan 2012: massive retro-orbital and medial orbital mass extending outwards through the medial orbital wall where a previous decompression with bone removal had facilitated spread of lesionsal tissue into the ethmoid sinus</td>
<td>Serial debubling, medial wall bony orbital decompression and dacryocystorhinostomy Exenteration with prosthetic reconstruction 2 courses chemotherapy for lung metastasis, stopped due to non-responsiveness of the metastases, patient intolerance, and poor prognosis</td>
<td>Vimentin+ SMA + Factor XIIIa + Calponin + Desmin - CD34 – For rhabdoid transformation: SMA + EMA + Calponin + Myogenin, myosin, myoglobin, muscle specific Actin, and Desmin were negative. IN1, GFAP, and S100 were also negative.</td>
<td>&gt;30 years seven recurrences</td>
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<td>Pujari et al 2018</td>
<td>Left conjunctiva</td>
<td>48/M</td>
<td>Conjunctival angiomyxoma</td>
<td>Conjunctival angiomyxoma</td>
<td>Painless fleshy conjunctival mass</td>
<td>B-scan ultrasonography: a well-defined mass without any intraocular extension. Contrast-enhanced computed tomography of the orbit revealed a well-defined, heterogeneously enhancing, mass lesion confined to the epibulbar surface without post-septal extension</td>
<td>Complete surgical excision with 2mm margin on inferior, nasal, and temporal side and up to superior fornix followed by amniotic membrane grafting</td>
<td>NA</td>
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SMA- Smooth Muscle Actin; EMA- Epithelial membrane antigen; GFAP- Glial fibrillary acidic protein; S-100- S-100 protein; CD- Cluster of Differentiation; INI1- Integrase Interactor 1

**Fig. 1:**

a) Clinical photograph of the patient at presentation.  
b) Clinical picture at 6 months postoperative follow up,  
c, d) Coronal and sagittal sections of CT scan Orbit, showing a heterogeneous lesion in the inferotemporal orbit indenting the left eyeball.
Fig. 2: a) Gross appearance of the lesion. b) Low power view to show tumour cells in a vascularized myxoid stroma. (H&E x200) c) High Power view shows bland spindle and stellate shaped cells. Inflammatory cells are also seen. (H&E x 400) d) Cytoplasmic Vimentin (Avidin Biotin x400) e) SMA positivity. (Avidin Biotin x400).
component to 63% in tumours with epithelial component [9]. Sites of predilection for superficial varieties involve
the trunk but can also appear on the lower limbs, head and
neck and pelvic region [4].

Patients are generally middle-aged adults, though cases
have been described in young children, usually presenting
with a painless gradually progressive mass which may
cause proptosis, diplopia, or diminution of vision [5].

Superficial angiomyxoma may form the earliest presenting
symptom in cases of Carney’s complex, especially
associated with external ears. Carney’s complex is an
autosomal dominant disorder comprising myxomas of the
heart and skin, hyperpigmentation of the skin (lentiginosis),
and endocrine overactivity [6].

Imaging by computed tomography or magnetic resonance
imaging is indicated which reveals a heterogeneously
enhancing locally aggressive mass with ill-defined margins
and infiltrating surrounding tissues without any bony
erosions or intracranial extensions. However, Hidayat et al.
described angiomyxomas in two children which manifested
locally aggressive behaviour with respiratory sinus,
cavernous sinus, and middle cranial fossa bone invasion [7].
This may underscore the fact that these tumours may have
a more aggressive course in children although distant
metastasis has not been reported on long follow-ups.

On gross examination, these tumours are irregular spongy
soft tumours which may contain mucoid material and are
well vascularized. The jelly-like consistency precludes
complete surgical excision. On microscopic examination,
these are low-cellularity tumours with bland stellate or
spindle-shaped cells in a loosely arranged myxoid matrix
that stains avidly with Alcian blue. Abundant vascularity
is a distinguishing feature from myxomas. One unique
feature, in contrast to other superficial myxoid lesions
such as digital myxoma or nerve sheath myxoma, is the
presence of neutrophils in the stroma associated with
necrosis or ulcer [8]. Thin-walled vessels are characteristic
of superficial angiomyxoma, whereas large-calibre vessels
with thicker walls are more classically found in the deep
aggressive variety. Excessive release of hyaluronic acid
from fibroblasts allows for infiltrative growth and proptosis
within tissues [9]. Some of these tumours have an epithelial
component associated with a higher rate of recurrence in
one study.

Immunohistochemical data pertaining to orbital
angioimmxomas is rather limited. In most cases, these
tumours have shown positivity for Vimentin, Calponin,
CD34 and Factor XIIIa. Negativity for SMA, S-100 and
Desmin has also been widely established [7].

Once a diagnosis is established, and any bony or intracranial
extension ruled out by imaging, complete surgical excision
is the treatment of choice. In many cases where this is not
possible, surgical debulking with a margin of safe tissue
should be undertaken. Chemotherapy and radiotherapy
have limited roles.

Distant metastasis has not been reported even on prolonged
periods of follow up. However, an isolated case of malignant
rhabdoid transformation with lung and other metastasis in
a longstanding case of aggressive angiomyxoma has been
reported which is labelled as a composite tumour, thus
necessitating early and extensive removal of the tumour [10].

To conclude, angiomyxoma is an extremely rare, locally
aggressive orbital tumor, occurring in the fourth to fifth
decade of life. The management of orbital angiomyxomas
can be quite challenging. A complete excision with margin
of healthy tissue has been recommended, but it is difficult
due to adhesions and the infiltrative nature of the tumor
and therefore tend to recur, requiring long-term follow-up.

Compliance with Ethical Standards
This report complies with ethical standards and permission
from the patient has been taken for publication

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Conflicts of interest
None

Written Informed Consent
The authors affirm that human research participant has
provided written informed consent regarding publishing
his data and photographs.

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