Malignant Orbital Meningioma Originating from the Frontal Lobe.

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MALIGNANT ORBITAL MENINGIOMA

ORIGINATING FROM THE FRONTAL LOBE

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Established Facts:

- Orbital meningiomas are uncommon and usually benign tumors, typically arising from the sphenoid bone or optic nerve sheath.

Novel Insights:

- An anaplastic orbital meningioma can originate from the frontal lobe of the brain, and can be associated with orbital and distant extracranial metastases.
- Orbital invasion may be more likely after surgical resection of aggressive frontal lobe meningiomas.

This Case was presented in part at the Verhoeff-Zimmerman Society meeting at Wills Eye Hospital, Philadelphia, Pennsylvania, April 22, 2017.
ABSTRACT

Orbital meningiomas are typically benign tumors, most commonly originating from the dura of the sphenoid wing or the optic nerve sheath. We describe an unusual case of a malignant meningioma originating from the frontal lobe that ultimately produced orbital and distant metastases. Orbital invasion by the meningioma was preceded by multiple incomplete resections, which may have facilitated access to the orbit. The present case serves to remind clinicians that surgical resection of aggressive, recurrent frontal lobe meningiomas may facilitate subsequent penetration of surrounding structures, particularly by tumors that demonstrate bone destructive properties.

KEYWORDS: meningioma, orbit, craniotomy, tumor recurrence, metastasis, exenteration
INTRODUCTION

Meningiomas are the most common intracranial neoplasms, typically occurring in the fourth through sixth decade of life [1]. They are usually benign, indolent tumors, twice as prevalent in women as men, with fewer than 10% demonstrating anaplastic features or distant metastases [1, 2].

Orbital meningiomas are estimated to account for 3-9% of all intraorbital neoplasms [3, 4]. Sphenoidal-origin meningiomas often narrow the optic canal and compress the optic nerve at that location. Primary orbital optic nerve meningiomas typically compress the nerve as they expand in the sub-dural space, and present with progressive, unilateral vision loss, axial proptosis, and optic atrophy.

Involvement of the anterior optic nerve may be associated with disc shunt vessels visible ophthalmoscopically. Clinical features of sphenoidal or other-origin intracranial meningiomas include, headaches, nausea and vomiting, and papilledema.

Primary orbital meningiomas arise from the optic nerve sheath meningothelial cells or rarely from ectopic arachnoid tissue, whereas secondary orbital meningiomas most commonly originate from the sphenoid wing [5]. We describe an unusual case of an orbital meningioma originating from the frontal lobe convexity, additionally remarkable for its aggressive behavior, anaplastic histology, multiple recurrences, local tissue destruction, and extracranial metastases.

CASE REPORT
A 54-year-old African-American woman was referred for an oculoplastic and orbital surgery consultation for orbital involvement by an anaplastic-subtype convexity meningioma of the right frontal lobe diagnosed 14 years prior. Initial recognition of the tumor had been preceded by recurrent headaches and seizures. Interval therapy had included four craniotomies, all demonstrating incomplete excision. Radiation therapy after the initial excision and following a subsequent excision were judged non-beneficial. Medical therapy had included levetiracetam and zolpidem.

Resection #2, 12 years after Resection #1, was performed with right frontal bone craniotomy and cranioplasty and demonstrated a 5.5 x 1 cm right frontal convexity dural based mass extending along the right skull base, without orbital or sinus involvement (Figure 1). Two additional resections were performed (#3 and #4) at intervals thereafter of 5 months and 14 months, respectively. Residual right frontal lobe tumor was noted on MRI after each surgery. Histopathology after each resection demonstrated a World Health Organization (WHO) grade III anaplastic meningioma with Ki-67 positive tumor cells, consistent with recurrence of the patient’s previously incompletely excised tumor. An MRI performed five months after Resection #4 demonstrated the first radiologic evidence of orbital involvement, revealing that the mass now extended through the roof of the right orbit into the superior extraconal space, although no intraocular involvement was evident (Figure 2A). The tumor displaced the globe inferiorly and temporally. The
patient then came under ophthalmic plastics care for further evaluation and
treatment of the orbital involvement.

At her plastics consultation the patient complained of right-sided orbital pain
and profound diminution in visual acuity in the right eye. She had no other past
ocular history or prior ophthalmic surgeries, and had no prior ophthalmic
examinations for comparison. On examination, a firm right upper lid mass was
present measuring 40 mm in the largest dimension. There was prominent proptosis
of the right globe and marked ptosis of the right eyelid (Figure 2B). Visual acuity
was light perception OD, which improved to 20/25 with manual elevation of the
right eyelid, and 20/20 OS. The right eye demonstrated a relative afferent pupillary
defect and the left pupillary reflex was normal. Intraocular pressures were 24 OD
and 18 mmHg OS. Confrontational visual field testing showed defects in all four
quadrants OD and no defects OS. Slit lamp examination of the anterior and
posterior segments was normal bilaterally. All MRIs were ordered by a
Neurosurgical consultant. After the initial MRI, a repeat 6 months later
demonstrated more extensive orbital invasion, prominent involvement of the right
frontal sinus, and loss of the cortical bony margin (Figure 2C). The tumor had
infiltrated the superior and medial rectus muscles, as well as the superior oblique
muscle and tendon and broached the orbital septum with subsequent progressive
lid involvement.

A right orbitotomy was performed to biopsy and characterize the orbital
mass. Tumor tissue was sampled from the superior nasal orbit, along with partial
resection of the superior rectus muscle. Intraoperatively, it was apparent that the
tumor had extensively infiltrated the soft tissue structures of the right orbit and, accordingly, total removal of the orbital mass could not be assured and was not attempted. Histopathology confirmed that the orbital mass was an anaplastic meningioma consistent with prior pathology.

Two months after the orbital biopsy the patient experienced intractable right eye and orbit pain poorly controlled by analgesics, worsening proptosis and orbital deformity, purulent discharge, persistent granulation tissue, and cutaneous erythema and scab formation (Figure 3). A right orbital exenteration was subsequently performed as a palliative measure to alleviate these symptoms.

Intraoperatively, exploration revealed that the tumor had eroded through the right frontal sinus and infiltrated the nasal two-thirds of the orbit displacing the globe inferiorly and temporally.

Pathologic examination of the exenteration specimen confirmed previous pathology and a diffusely infiltrating tumor of the orbit and eyelids. Tumor cells were pleomorphic with large vesicular nuclei, abundant eosinophilic cytoplasm, and frequent mitoses (Figure 4). Immunohistochemical staining of the orbital tumor was positive for CD68, epithelial membrane antigen (EMA), moderately high proliferation index (Ki-67), progesterone receptor (PR), and somatostatin receptor type 2 (SSRT-2). Tumor was present in the conjunctival stroma and eyelid dermis. Sections of the globe demonstrated reduction in retinal ganglion cells and early optic atrophy but no scleral or intraocular tumor.

Two months after the exenteration, the patient began experiencing dyspnea, hemoptysis, and chest pain. CT of the chest, abdomen, and pelvis demonstrated
multiple well-defined intrapulmonary nodules bilaterally. A CT core biopsy of the left lung confirmed metastatic anaplastic meningioma. (Figure 4). External iliac lymph nodes were also prominent in imaging but not biopsied. The patient’s clinical condition subsequently deteriorated and she died from respiratory impairment five months later while in hospice care.

**DISCUSSION**

The present case demonstrates a secondary orbital malignant meningioma originating from a highly unusual primary site, the frontal lobe convexity. The mechanism of orbital invasion in this case is uncertain, although the patient’s history of multiple craniotomies preceding the first radiographic evidence of orbital involvement could have played a significant role. The patient underwent a total of four frontal bone craniotomies before orbital involvement became apparent. Highly malignant intracranial tumors such as glioblastoma multiforme have been observed to infiltrate the orbit through a prior craniotomy site [9-11], although, to our knowledge, invasion of the orbit by a meningioma through a craniotomy site has not been proven. The possibility of hematogenous spread from the frontal convexity to the orbit is less likely in this case, given the presence of only a single mass in the orbit rather than multiple distinct foci, and strong imaging evidence of direct extension.

Alternatively, the tumor may have penetrated the orbit by gross destruction of orbital bone. Malignant meningioma subtypes may exhibit bony destruction or
hyperostosis [12]. Sphenoid wing meningiomas, in particular, demonstrate a high incidence of bone involvement [13], and cases of intraosseous meningiomas can display local bone destruction. Extension into the orbit through bone has also been previously reported in cases of glioblastoma multiforme, and, in rare cases, by pituitary tumors and craniopharyngiomas [14]. Interestingly, various reports describe the phenomenon of convexity meningiomas eroding through bone in the absence of prior craniotomies, though not into the orbit specifically [15, 16].

An additional peculiarity of this case is the highly malignant histology and aggressive behavior of the meningioma, with multiple recurrences likely due to incomplete excision, and eventual metastasis to the lungs and iliac nodes. Meningiomas arising over the convexities of the brain are generally benign, and typically have high potential for complete surgical excision, if a substantial margin of excision can be achieved. However, those with anaplastic pathology, as in the present case, can demonstrate 5-year recurrence rates as high as 50% [17].

Meningiomas are rarely metastatic, with distant extracranial metastases estimated to occur in approximately 0.001% of cases, most commonly to the lung [23]. As expected, the tumor described in this case exhibited characteristics associated with a higher risk for metastasis, including histologic malignancy (WHO grade III), and local recurrences. Adlakha et al. in 1999 suggested that previous craniotomy may be a predisposing factor for metastasis, although the mechanism underlying this association has not been clearly elucidated [18].

In conclusion, this case illustrates a secondary orbital meningioma arising from an unusual site of origin, the frontal lobe. Though the mechanism of orbital
invasion in this case is uncertain, clinicians should be aware that a history of multiple craniotomies to resect aggressive frontal lobe meningiomas may confer a higher risk of both orbital involvement and subsequent metastases.

FIGURES

Figure 1. Coronal (A) and sagittal (B) enhanced CT scans showing an entirely intracranial dural based mass in the right frontal pole with no extension into the orbital roof or frontal bone. There is no involvement of the sinuses or the falx. There is considerable surrounding edema in the frontal lobe.
Figure 2. (A) Coronal MRI scan (T1 weighted following gadolinium contrast infusion) performed after four tumor resection surgeries. There is a lobulated enhancing extraconal tumor mass located in the right superior orbit compressing the globe inferiorly and temporally. No intraocular invasion is evident. (B) Facial photograph of patient demonstrating a firm right upper lid mass measuring 40 mm in the largest dimension. There is prominent right-sided proptosis and near complete ptosis. (C) Axial MRI scan (T1 weighted fat-saturated images following gadolinium contrast infusion) performed six months after the scan depicted in (A). The enhancing combined intra- (arrow) and extra-cranial tumor mass is well seen. The extra-cranial component extends into the superior medial portion of the orbit and displaces the globe temporally. There is tumor in the intervening frontal sinus as well with loss of the cortical bony margin.
Figure 3. Facial photograph of patient two months after a subtotal tumor resection was performed, demonstrating worsening proptosis and orbital deformity, purulent discharge, persistent granulation tissue, and cutaneous erythema and scab formation.
Figure 4. (A) Photomicrograph of orbital extension of frontal lobe malignant meningioma demonstrating margin between tumor tissue that is viable (lower right side) and necrotic (upper left side, labeled). (H&E original magnification X 125). (B) Higher magnification photomicrograph of orbital meningioma demonstrating pleomorphic tumor cells with large vesicular nuclei and prominent nucleoli, including two abnormal mitoses (arrows). (H&E original magnification X 500). (C) Photomicrograph of IHC Somatostatin-2 which is highly positive in the orbital lesion and consistent with meningioma. (SSRT-2 original magnification X 125). (D) Photomicrograph of lung metastases biopsy (CT Core sample) from left lung including the margin of pulmonary tissue (PT) on the right. The inset at higher magnification demonstrates an area of the same
biopsy with histologic characteristics similar to those in the orbital lesion. (H&E original magnification X 125; Inset: H&E original magnification X 500)

REFERENCES


