Lateral and Middle Sphenoid Wing Meningioma

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Approximately 15 to 20% of all meningiomas arise from the sphenoid wing, with half of them emanating from the lateral and middle portion of this bone. Lateral sphenoid wing meningiomas arise from the pterion and typically grow along the Sylvian fissure.

There are no pathologic or genetic features specific to lateral and middle sphenoid wing meningiomas, although World Health Organization (WHO) class II and III meningiomas are less common in this region of the brain than over the convexities.

Classification

Sphenoid wing meningiomas can be divided into three main groups: 1) meningiomas arising from the anterior clinoid and medial third of the sphenoid wing, 2) meningiomas arising from the middle and lateral sphenoid wing, and 3) en plaque meningiomas of the sphenoid wing or spheno-orbital meningiomas.

In my opinion, surgical classification should simplify stratification of surgical objectives. I therefore divide lateral and middle sphenoid wing meningiomas into two main categories: 1) globular or traditional tumors, and 2) hyperostotic en plaque or spheno-orbital tumors.

Globular tumors are relatively straightforward and can be transformed into convexity tumors after resection of the lateral sphenoid wing. However, spheno-orbital tumors are more technically challenging because expansile osteotomy is necessary to remove a reasonable
portion of the tumor-infiltrated bone and correct the proptosis. Intraorbital extension of these tumors is not uncommon.

In this chapter, I discuss meningiomas of the middle and lateral portions of the sphenoid wing. Large tumors of the middle portion of the sphenoid wing involving the optic apparatus, superior orbital fissure, or cavernous sinus are discussed in the Medial Sphenoid Wing Meningioma Chapter. Spheno-orbital meningiomas of the entire breadth of the sphenoid wing including those affecting the anterior clinoid process are reviewed in this offering.

**Diagnosis**

Lateral and middle sphenoid wing meningiomas typically present when they have reached a large size. Patients may present with headaches, contralateral hemiparesis, focal motor seizures, or aphasia (tumors on the dominant hemisphere). Hyperostosis of the sphenoid wing may cause exophthalmos, orbital pain, blurry vision, diplopia, or visual field dysfunction. Hyperostosis of the lateral aspect of the sphenoid wing or adjacent calvarium may also cause cosmetic concerns such as a bump on the head.

Large tumors causing temporal lobe edema may result in focal, complex-partial, or generalized tonic-clonic seizures including gustatory and olfactory hallucinations. Neuropsychological disturbances often exist, but are apparent only to patients and their families in retrospect. These lesions may also be found during diagnostic tests for headaches, vertigo, or other unrelated complaints.

**Evaluation**

Thin-cut magnetic resonance (MR) images including the fat suppression sequences through the orbits assess the extent of orbital involvement. Computed tomography (CT) with bone windows and thin slices measures the extent of bone invasion or hyperostosis.
Consultation with our oculoplastic colleagues is very helpful for baseline evaluation of vision and proptosis. Preoperative preparation of a custom implant to reconstruct the walls of the orbit and lateral calvarium after resection of large areas of affected bone may be needed.

The T2-weighted MR images are also essential for identifying vasculature and intraparenchymal edema. Preoperative embolization is usually not helpful for lateral sphenoid wing meningiomas because the craniotomy typically allows for devascularization of the blood supply (primarily from the middle meningeal artery) early during surgery.
Figure 1: A classic left-sided lateral sphenoid wing meningioma with sphenoid wing hyperostosis (top images) is shown. A right-sided spheno-orbital meningioma (bottom images) with significant hyperostosis of the sphenoid wing, including the clinoid and orbital walls, causing proptosis, is also demonstrated. The resection of both the intradural tumor and hyperostotic bone is necessary for achieving long-term tumor control and correction of the proptosis.
**Indications for Procedure**

Lateral sphenoid wing hyperostosis and tumor extension into the orbit leads to proptosis and vision deterioration. These latter symptoms are clear indications for operative intervention. Hyperostosis may also cause optic nerve compression at its foramen. As is the case with other meningiomas, mass effect, intracerebral edema, and associated symptoms such as seizures, cognitive impairment, and speech difficulties are other indications for intervention.

![Image of lateral sphenoid wing meningioma](image)

**Figure 2:** A right-sided lateral sphenoid wing meningioma and its relationship to the surrounding structures is illustrated.

**Preoperative Considerations**
Surgical resection is the mainstay of treatment for lateral sphenoid wing meningiomas, and the goal of surgery is complete removal of the tumor with wide excision of the affected bone and dura, minimizing the risk for future tumor recurrence.

Large and multicompartmental tumors with involvement of the superior orbital fissure or cavernous sinus may not be curable in this manner. Stereotactic radiosurgery may be employed for small residual or recurrent growing tumors, especially for portions of the tumor invading the cavernous sinus.

As mentioned previously, a custom cranial implant is prepared preoperatively to minimize the risk of significant postoperative cosmetic deformity after resection of a large portion of the calvarium and orbit affected by the hyperostosing tumor.

I prefer the use of a lumbar drain for dural decompression and unobstructed extradural osteotomy. Cerebrospinal fluid (CSF) drainage facilitates dural and brain mobilization and prevents their injury during the drilling process.

**Operative Anatomy**

The anatomy of the sphenoid bone is briefly described below.
Figure 3: The anatomy of the sphenoid wing and clinoid process in the coronal planes as it relates to lateral and middle sphenoid wing meningiomas (images courtesy of AL Rhoton, Jr).
Figure 4: The anatomy of soft tissues around the sphenoid bone relevant to removal of sphenoo-orbital meningiomas. Note the location of the frontal nerve, levator, and superior rectus muscles. Many lateral and middle sphenoid wing meningiomas invade the superior orbital fissure and preclude their gross total resection (images courtesy of AL Rhoton, Jr).

RESECTION OF LATERAL AND MIDDLE SPHENOID WING MENINGIOMA

The patient is placed in the supine position with his or her head turned and immobilized in a skull clamp approximately 40 degrees away from the side of the approach. The use of prophylactic perioperative anticonvulsant medications is controversial, but I prefer to give a loading dose prior to surgery and continue the medication for 7 days postoperatively if pial invasion by the tumor is evident.

If the patient demonstrates any seizure activity preoperatively or during the postoperative time, the dose can be increased, and the
duration lengthened to 6 month to 1 year. Tumors with no or minimal intradural extension do not require the use of anticonvulsant medications.

Lateral and middle sphenoid wing meningiomas are resected through the pterional craniotomy. Neuronavigation (using MR and CT) guides the size of the craniotomy based on the extent of dural infiltration by the tumor. A sincere attempt should be made to resect the tumor’s dural tails. Since the dura is heavily infiltrated in these tumors, a generous piece of pericranium is harvested beyond the scalp edges during exposure and used to reconstruct the defect in the dura at the end of the procedure.

I also use a lumbar drain intraoperatively to assist with brain decompression and cortical dissection. Since the basal cisterns are not accessed until the last steps of the procedure, CSF drainage through the lumbar drain significantly assists with brain relaxation, making handling of the tumor and extradural drilling of the bone (especially in the case of spheno-orbital tumors) safe while avoiding frontotemporal lobar retraction.
Figure 5: Following reflection of the myocutaneous flap, I use a large burr to drill the hyperostotic calvarium that is centered over the pterion; a craniotomy is not performed initially (top image, right-sided approach corresponding to the lower images in Figure 1). Alternatively, multiple burr holes may be placed at the periphery of the tumor-infiltrated bone; these holes are then connected using the craniotome (bottom image). The hyperostotic bone is highly vascular and its elevation in one piece will often lead to excessive bleeding. In addition, the tumor-infiltrated bone and dura are adherent to each other, and elevation of the bone flap in one piece leads to avulsion injury to the cortex.

Figure 6: As each burr hole is created in the periphery of the tumor-infiltrated calvarium, bone wax is used to control bleeding from the thick cancellous bone (left image). Rongeurs may be used to remove the thickest portion of the bone at the pterion and the lateral aspect of the sphenoid wing (right image). The enlarged middle meningeal arteries feeding the tumor should be coagulated.
Figure 7: After the pterion and the affected island of bone are drilled away, the normal edges of the bone and dura are exposed. Next, an orbitozygomatic craniotomy is completed (top image), and the affected medial sphenoid wing (*) is isolated using a #1 Penfield dissector (bottom image). This bone is drilled away using a small side-cutting burr.
Figure 8: Aggressive elevation of the orbitozygomatic or pterional bone flap without its complete disconnection under direct vision along the orbital roof or lateral sphenoid wing should be avoided. Attempts to elevate the bone flap for the tumor in the top images in Figure 1 led to an inadvertent extension of the fracture along the sphenoid wing into the optic canal (top image) and ethmoid sinus (bottom image) with resultant optic nerve injury and unrecognized postoperative CSF leak.
Figure 9: After removal of the medial sphenoid wing and extradural clinoidectomy, the superior orbital fissure (SOF) and optic nerve (ON) are identified. At this stage, bony resection is guided by intraoperative CT navigation. If the tumor extends into the orbit, the superior and lateral walls of the orbit are removed further and the intraorbital extension of the tumor is excised. An implant is needed to reconstruct part of the orbital walls at the end of the procedure to avoid delayed endophthalmus.

INTRADURAL PROCEDURE

Removal of the intradural component(s) of the tumor is conducted using the same techniques as those for convexity meningiomas.
Figure 10: The dura is then incised sharply while the operator protects the brain, and the dural edges are coagulated using bipolar electrocautery. If the dural tails of the tumor extend beyond the initial bony exposure, the craniotomy is expanded to ensure the infiltrated dura has been generously exposed and resected. MR navigation can guide the operator regarding the extent of dural resection. The intradural component of this sphenoid-orbital meningioma reached into the anterior Sylvian fissure.

The intradural tumor is devascularized along its dural base. Dissection of the tumor from the surrounding brain is carried out circumferentially, while the operator places cottonoid patties at the periphery of the tumor and mobilizes the decompressed tumor away from the brain.

At this point, it is important to identify and preserve the distal branches (M3 and M4) of the middle cerebral artery. These branches, along with the main middle cerebral artery trunk, may be encased in the tumor, but usually a dividing arachnoid plane is present and there is no invasion of the arterial wall.
Once the tumor is completely dissected and removed, any remaining nodules or tumor-infiltrated dura are excised. Most of the affected dura is resected, however, the affected dura at the level of the superior orbital fissure and the cavernous sinus should be left intact.

**Additional Considerations**

Some meningiomas may infiltrate the dura and bone extensively.

Figure 11: If the tumor has dramatically invaded the cortex, dura, and bone, the latter two layers may have to be removed *en bloc*. To do so, the bone is not elevated until the dura is cut along the bony margins and the underlying attached tumor dissected away from the brain. This maneuver is appropriate for small and superficial tumors.
Figure 12: The tumor is dissected away as the operator uses cottonoid patties to “wipe the brain away” from the tumor (inset image).
Figure 13: Gross total resection is possible through elevation of all of the tumor-infiltrated tissues.

**Closure**

The pericranial graft harvested at the beginning of the procedure is placed over the dural defect and sewn in place.
Figure 14: A custom implant is used to reconstruct the orbital rim and part of the orbital walls without compromising the movement of eye muscles. Cranioplasty using titanium mesh repairs the remaining defect in the calvarium. The tumor-infiltrated bone flap is not replaced.

Pearls and Pitfalls

- A lumbar drain assists with brain relaxation during extradural bone work for resection of an sphenoid-orbital meningioma. The arachnoid cisterns are not accessed until the later stages of the operation for a lateral pterional meningioma.
- The hyperostotic lateral sphenoid wing, pterion, and calverium can be highly vascular, and excessive bleeding may be encountered during bone work. Therefore, placement of burr holes at the margins of the affected bone and stepwise hemostasis using bone wax can minimize blood loss.
- Aggressive elevation of the orbitozygomatic or pterional bone
flap without its complete disconnection under direct vision along the orbital roof or lateral sphenoid wing should be avoided. Attempts to elevate the bone flap along the hyperostotic lateral sphenoid wing can lead to an inadvertent extension of the fracture into the ethmoid sinus and optic canal with resultant unrecognized postoperative CSF leak or optic nerve injury.

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References


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<table>
<thead>
<tr>
<th>Title</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
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</tr>
<tr>
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</tr>
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</tr>
<tr>
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</tr>
<tr>
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</tr>
<tr>
<td>Management of the optic canal invasion and visual outcome in sphe...</td>
</tr>
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