Surgical treatment of Spheno-Orbital Meningiomas
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Surgical treatment of sphenoorbital meningioma

Competing interest: None to declare.

Abstract

Purpose: To evaluate the outcome of surgery and radiotherapy in the treatment of sphenoorbital meningioma (SOM).

Method: A retrospective study of 66 consecutive cases treated with surgery for SOM with a minimum follow-up of 4 years. Clinical and radiological information were compared before and after the following surgical approaches: frontotemporal craniotomy, frontotemporal craniotomy combined with orbitozygomatic resection, and extended lateral orbitotomy alone.

Results: The median age at presentation was 46 (range, 26–68) years and the median follow-up after surgery was 102 (48–288) months. In total, 48 (73%) patients showed preoperative visual deterioration, with visual field defects. All patients had proptosis at presentation (mean±SD = 6.4±3.0 mm). Surgery for patients with SOM arrested visual deterioration in 61% and improved vision in 30% of cases. Furthermore, a substantial reduction of proptosis was achieved in 85% of patients. The proptosis in our group was reduced by 2.6±2.6 mm. There was no correlation between surgical approach and proptosis reduction ($P = 0.125$). The recurrence rate was 17%. Only one of 15 patients who underwent radiotherapy showed signs of recurrence.
**Conclusions:** The surgical aims in the treatment of SOM should be the restoration of visual acuity and reduction of proptosis, rather than complete tumour removal. The surgical approach can be tailored to individual cases. We recommend radiotherapy in cases of subtotally removed SOM.

**Introduction**

Sphenoorbital meningiomas (SOM) originate from the dura of the sphenoid wing, involve the orbit and cause visual deterioration and proptosis.\(^ \text{1-12} \) Given their extensive dural and orbital involvement, adequate resection of SOM is difficult, leading to a recurrence rate of up to 50%.\(^ \text{13-15} \) The main aim of surgery is to restore the deteriorated vision and treat proptosis.\(^ \text{1-12} \) Adequate treatment strategies and surgical approaches remain controversial. Some authors have reported that longstanding tumour-related proptosis cannot be treated surgically and that it should therefore not be attempted.\(^ \text{16} \) Others advocate early, aggressive surgical therapy as the initial treatment.\(^ \text{17} \) Postoperative radiotherapy is recommended for subtotally removed or recurrent SOMs,\(^ \text{18} \) although little information is available regarding the efficacy and side effects of this treatment.

This study was performed to evaluate different surgical approaches used in two centres over a period of two decades. We studied long-term surgical outcomes and the effects and complications associated with postoperative radiotherapy in SOM.

**Patients and Methods**

We examined the records of 90 patients treated between 1980 and 2007 at the Orbital Centre of the Academic Medical Centre, University of Amsterdam (\( n = 61 \)), and at the
University Medical Centre in Utrecht \((n = 29)\) with a diagnosis of SOM. Sixty-six patients in this group underwent surgery at the Orbital Centre of the Academic Medical Centre, University of Amsterdam \((n = 40)\), and the University Medical Centre in Utrecht \((n = 26)\). Fifty-one patients with optic neuropathy combined with disfiguring proptosis and 15 with disfiguring proptosis were indicated for surgery. Twenty-three patients underwent no surgery due to stable visual acuity and the absence of optic neuropathy. Despite visual deterioration in the amblyopic eye, one patient declined surgical intervention. Only patients who underwent surgery were included in this study.

The clinical history, surgical approach, and outcome were obtained retrospectively from a review of patient charts and imaging. Our analysis considered neurological symptoms and their duration, ophthalmological examinations, consisting of testing the patient’s visual acuity (Snellen notation) and pupil responses, funduscopy, tonometry, and Goldmann or Humphrey perimetry for visual field defects. Ocular motility was also evaluated.

Diplopia was considered present when the patient experienced double vision (in any gaze direction) that disappeared with one eye closed. The degree of proptosis was determined using a Hertel exophthalmometer. In Amsterdam, an Oculus Hertel (Oculus Optikgeräte GmbH, Wetzlar, Germany) was used and in Utrecht, a Zeiss Hertel (Carl Zeiss, Jena, Germany) was used.

All patients underwent preoperative computed tomography (CT) or magnetic resonance imaging (MRI; \(n = 51\)) with contrast. Imaging was scheduled immediately after surgery, at 3 and 12 months postoperatively, and annually thereafter for 5 years, and then at longer intervals, depending on the presence or absence of residual or recurrent tumours.
In patients who underwent radiotherapy, tear production was assessed using the Schirmer test and corneal dryness was classified as no corneal change, corneal stippling, limited to the inferior periphery, more extended stippling, ulceration and clouding postradiation cataract. When necessary, fluorescein angiography was performed to evaluate any postradiation retinopathy. After radiotherapy, all patients underwent detailed endocrinological testing.

Surgical procedures

Frontotemporal (pterional) craniotomy was performed as described. The greater wing of the sphenoid ridge was removed to the lateral limit of the lesser wing. When hyperostotic, the anterior clinoid was removed extradurally and the superior orbital fissure (SOF) was unroofed. The optic canal was unroofed, either intra- or extradurally, and the inferior optic strut was removed. When intracranial or intraorbital soft tissue was present, it was resected.

Orbitozygomatic craniotomy (OZ) is an extension of the frontotemporal approach. The superior and lateral orbital rims were mobilised with additional removal of part of the lateral wall along the zygoma and orbital roof, which provides access to the floor of the anterior and middle cranial fossae. This was performed in either one or two pieces, with the superior and lateral rim.

Extended lateral orbitotomy was performed through either a coronal or lid crease incision. In this approach, the classical lateral orbitotomy bone incision described by Berke was enlarged from the deep sphenoid wing up to the SOF.
Reconstruction

Periorbital reconstruction was not performed. The superior and lateral orbital rim bone was reattached. The roof and lateral wall of the orbit were routinely reconstructed with a split cranial bone graft and secured to the orbital rim by titanium miniplates. In two cases, this was performed using titanium mesh. Dural defects were reconstructed with autologous galea grafts, which included convexity and basal defects. If there was a large cavity between the orbital and dural reconstruction and the cranioplasty, an autologous fat graft, harvested from the abdomen, was used to fill this and prevent a postoperative pseudomeningocele. The temporal muscle was resuspended and sutured meticulously.

Radiotherapy

Postoperatively, 15 patients underwent radiotherapy of 54 Gy in 1.8-Gy fractions. The inclusion criteria for radiotherapy were large rest tumour and involvement of the cavernous sinus, while the exclusion criteria were the presence of diabetes mellitus and hypertension.

Statistical Analyses

Visual acuity and pre- and postoperative proptosis are described as means±SD. Differences between the three different surgical approaches were compared using the Kruskal–Wallis test. Proptosis reduction was compared between the three approaches with correction for preoperative proptosis (nonparametric ANOVA). All analyses were carried out using the SPSS software (ver. 16 for Windows; SPSS, Chicago, IL, USA). In all analyses, $P < 0.05$ was deemed to indicate statistical significance.
**Results**

The median age at presentation was 46 (range, 26–68) years. The study population consisted of 61 (92%) females and five (8%) males. The left and right orbits were involved in 35 and 31 patients, respectively. The median follow-up after surgery was 102 months (48–288 months) and the median period between presentation and surgery was 24 months (1–119 months). In total, 51 (77%) patients had preoperative progressive visual deterioration with visual field defects. Three patients had no light perception (NLP) preoperatively. All patients had proptosis at presentation, with a mean of 5.89±2.89 mm (mean±SD). At presentation, seven (11%) patients had diplopia and 19 (29%) had headaches or retrobulbar pain.

**Surgery**

The surgical approach was frontotemporal craniotomy in 45 patients and this approach was combined with orbitozygomatic resection in 10 patients. Forty-six patients in this group had optic neuropathy due to compression of the optic nerve, five had disfiguring proptosis with hyperostosis of the optic canal and four had proptosis combined with retrobulbar pain. As all of these 55 patients had disfiguring proptosis, this was also addressed at the time of surgery.

An extended lateral orbitotomy was performed in 11 patients. Patients in this group had only disfiguring proptosis and two had severe proptosis with stretching of the optic nerve.
Craniotomies

Fifty-five patients underwent a large unilateral frontotemporal craniotomy, followed by removal of additional tumour-infiltrated bone via an extradural approach in all cases. Ten procedures also included OZ resection. The orbital roof was resected in 49 cases and the lateral orbital wall was removed in 35. The optic canal and SOF were opened and decompressed in 51 and 29 cases, respectively. After opening the orbit, periorbital resection or stripping was necessary in 32 procedures. Extensive en plaque dural involvement was found in 45 cases; resection of the convexity-dura was necessary in 39 cases.

Reconstruction was performed in 9 of 10 patients treated via the frontotemporal approach in combination with OZ. The reconstruction used calvarial bone in seven patients and titanium mesh in two; both the lateral and orbital roof were reconstructed in all nine patients (Fig. 1).

Orbitotomy

In 11 patients who underwent lateral orbitotomy, the lateral wall of the orbit until the SOF was removed; in five cases, the orbital roof was also partially removed. In six procedures, periorbital resection was also performed due to tumour infiltration. In five cases in which there was infiltration of the ocular muscles, only the exophytic tumour was removed. None of these patients underwent reconstruction. All lateral orbitotomies were performed by orbital surgeons (Fig. 1).

Radiotherapy

Fifteen patients underwent secondary radiotherapy after surgical excision. All 15 had
significant residual tumours, which also involved the cavernous sinus in four cases. Two
patients underwent radiotherapy after debulking of a recurrent lesion. Only one of these
patients showed further recurrence after follow-up for 38 months. In four patients, the
tumour volume was reduced to less than 10%. In all four, there was significant soft
tissue involvement.

**Outcome**

**Visual acuity**

Forty (61%) patients showed either stabilisation of their visual acuity or an
improvement of less than two lines on the Snellen chart. Three eyes from this group
maintained a visual acuity of NLP (no light perception). Fifteen had a visual acuity of
more than 0.8 preoperatively, which remained stable postoperatively. Twenty (30%) had
improved visual acuity. Eight improved by more than four lines on the Snellen chart and
12 others by more than two lines.

There was no significant difference ($P = 0.195$) in postoperative vision between
the different approaches. All except two patients with visual deterioration underwent
decompression of the optic canal through a craniotomy. The two exceptions had
proptosis of more than 10 mm with stretching of the optic nerve and underwent only
extended lateral orbitotomy with total removal of the lateral orbital wall.

**Proptosis**

The mean proptosis, measured with a Hertel exophthalmometer, in this group was
6.4±3.0 mm preoperatively and 3.8±2.3 mm postoperatively. The mean reduction in
proptosis was 2.6±2.6 mm. Seventeen orbits showed a proptosis reduction of more than
5 mm, while ten showed no change in proptosis compared with preoperative values. One patient who underwent no reconstruction of the orbit developed enophthalmos of 2 mm. Fifty patients had a residual proptosis exceeding 2 mm. Patients who underwent lateral orbitotomy and combined craniotomy and OZ had the greatest mean preoperative proptosis of 7.9±3.4 and 7.7±2.9 mm, respectively. The mean proptosis with frontotemporal craniotomy only was 5.9±2.9 mm. However, there was no significant ($P = 0.125$) difference in the postoperative reduction in proptosis among the approaches (Fig. 2).

**Diplopia**

New diplopia was observed in 40 patients; it was transient in 32 cases. Two patients showed no eye movement in the first weeks after surgery; in both patients, the lateral wall was removed completely. Within 6 months, eye movement recovered completely, with no diplopia. Eight patients (12%) still showed diplopia; three of these underwent strabismus surgery with satisfactory results. In six patients after a frontotemporal craniotomy, the cause of the diplopia was third nerve palsy; thus, no strabismus surgery was performed. Ptosis surgery was performed in one patient with partial third nerve palsy and no diplopia. Nine patients with diplopia in the primary position preoperatively still had diplopia postoperatively and underwent surgical correction.

**Pain**

Of 19 patients with retrobulbar pain after surgery, 16 had no further pain. All of these patients underwent craniotomies.
Complications

Surgery

Five patients developed subgaleal cerebrospinal fluid (CSF) accumulation, which regressed spontaneously in four cases; one patient required surgical revision of the dural leak and placement of a lumboperitoneal shunt. After craniotomies, permanent third nerve palsies were seen in six cases, and fourth and sixth nerve palsies occurred in one patient each. Permanent trigeminal and facial nerve palsies were present in six and three patients, respectively.

In two patients who underwent OZ, the lateral excursion of the mandible was reduced by approximately 50%.

Radiotherapy

Three patients developed severe punctate keratopathy necessitating the daily use of lubricants. One patient also developed radiation retinopathy, but had stable visual acuity of 0.5 after laser treatment. During follow-up, one patient developed pituitary function impairment. There was no death due to treatment or SOM in this study population.

Recurrence

Eleven (17%) patients showed recurrence (regrowth). The median time between surgery and recurrence was 46 months (10–108 months). Macroscopic radical excision was achieved in only six patients. Of 61 patients with rest tumour, 15 underwent radiotherapy. Despite the larger size of the rest tumour in this group, only one of these 15 patients (7%) showed recurrence.
The signs of recurrence in four patients were a combination of visual
deterioration and increased proptosis, while four patients showed only a mild increase in
proptosis (< 3 mm). Four patients with visual deterioration underwent second
craniotomy—two had improved visual acuity and two other patients with NLP
underwent two other debulking surgeries and the affected eye was ultimately enucleated
because of corneal ulceration caused by proptosis.

Discussion
Symptom-oriented surgery for SOM is directed primarily at optic nerve decompression
when there is decreased visual acuity. In our series, the visual acuity improved in 30%
of cases and remained the same in 61% of cases. The cause of optic neuropathy was
addressed and treated in all of the patients with optic neuropathy. This was probably
related to the degree and period of optic nerve compression. In our series, the proptosis
was reduced by a mean of 2.6 mm from preoperative values, ranging from 2 to 14 mm.
Seventeen (26%) patients showed a marked reduction in proptosis by more than 5 mm
and up to 11 mm. Excluding one case of enophthalmos, residual proptosis was present
in 50 (76%) of the remaining patients. Nevertheless, 85% showed an improvement of
more than 2 mm. All patients in our series underwent periodical proptosis measurements
with a Hertel exophthalmometer. There have been only a few previous attempts to
quantify postoperative proptosis in SOM.\textsuperscript{2,5,12} Ringel \textit{et al.} reported improvement of
proptosis in 43 of 56 patients (77%) with no change in proptosis in the remaining 13
patients (23%).\textsuperscript{2} A recent study quantified the proptosis by MRI and reported significant
residual proptosis in 53% of cases.\textsuperscript{5} We believe that the cause of this significant residual
proptosis was long-standing congestion and fibrosis, resulting in less mobile orbital
contents. Despite this residual proptosis in the surgical treatment of SOM, the proptosis improves in most cases. Surgery also seemed to be effective at relieving pain associated with SOM; 16 of the 19 patients with reported pain showed no sign of pain postoperatively. We believe that pain is an underreported symptom in these series.

Our recurrence rate of 17% was consistent with other recently published series with long-term follow-up.\textsuperscript{1-12} These recurrence rates are much lower than those in earlier studies, which reported recurrence rates of 30–50%.\textsuperscript{13-14} Maroon \textit{et al.} listed the following reasons for the high recurrence rate in SOM: failure to diagnose the tumour early because the symptoms were confused with those of other clinical entities, such as fibrous dysplasia and Graves’ disease; inadequate resection, due to the involvement of important neurovascular structures; and the surgeon’s concern of iatrogenic death and serious complications in radical resections.\textsuperscript{18}

New imaging techniques appear to have improved the early diagnosis and effective surgical resection. To resect SOM, different surgical approaches are used, including transzygomatic, pterional, frontotemporal, and combined transcranial-transmalar and cranioorbital approaches. All of these approaches allow sufficient access to the orbit and middle fossa base for bony and soft-tissue tumour resection and decompression of the SOF and optic canal.\textsuperscript{19,20,23,24} Extended lateral orbitotomy or total lateral orbitotomy has been used routinely for severe cases of Graves’ orbitopathy and tumours in the orbital apex. This approach is effective in patients with SOM in which the main symptom is disfiguring proptosis, without involving the risks of a craniotomy.\textsuperscript{21,22,25,26} Recently, Lund reported 12 patients with SOM who underwent endoscopic endonasal medial orbital wall decompression and decompression of the optic canal in eight patients with optic neuropathy and visual deterioration.\textsuperscript{27} In these
patients, the opticopathy improved and the visual acuity improved by 1–4 lines on the Snellen chart.

Depending on the extent of the orbital wall and roof resection, most authors recommend firm reconstruction of the orbital walls to avoid pulsating enophthalmos and diplopia.

Maroon et al. reported 200 cases of orbital wall and roof resection without reconstruction, in which no cases showed permanent pulsating enophthalmos.18 DeMonte et al. concluded that partial or complete orbital roof resection, isolated or combined with lateral or medial orbital wall defects, did not require routine reconstruction when the periorbita was not resected.28 We advocate reconstruction of the orbital roof and the lateral orbital rim. We consider reconstruction of the lateral orbital wall to be unnecessary in most cases. Studies of orbital decompression in Graves’ orbitopathy have shown that lateral wall decompression can reduce the proptosis by up to 2 mm.29,30 As significant residual proptosis was present with all surgical approaches in SOM in these series, no further reconstruction of the lateral orbital wall may contribute to further proptosis reduction, while reconstruction of the orbit roof is sufficient to avoid pulsating enophthalmos. Techniques have improved since 1952, when Castellano et al. reported a high surgical mortality rate (20%) associated with SOM and concluded that SOM should be used only as a last resort.16 Although the mortality rate has decreased, some recent studies still reported mortality rates of 3–6%.2,3,11,26 Other complications, such as vision loss and transient (6–84%) or permanent (7–30%) cranial nerve deficits, are frequent.1–12 Although life-threatening complications were not seen in this series, postoperative visual deterioration occurred in 6 of 66 cases (9%).
oculomotor nerve palsy in 5 of 66 cases (7%) patients, and facial nerve palsy in 3 of 66 cases (5%).

Radiotherapy is often used to treat SOM. Peele et al. delivered a total dose of 45 Gy to subtotally or recurrent SOMs and none of the 42 patients in the radiation treatment group experienced recurrence during an observation period of 4.2 years, whereas 21 of the 44 patients in the non-radiation treatment group later presented with recurrent tumour growth.31

Another study indicated a decrease in size of skull base meningiomas in 51% of patients, stabilisation in 47% of cases and increased tumour size in 2% of cases.32 In our series, postoperative radiotherapy in subtotally removed SOM resulted in a reduction in tumour size and stabilisation in 27% and 67% of cases, respectively.

In conclusion, the aim of surgery in the treatment of SOM should be restoration of visual acuity and reduction of proptosis, rather than complete bony tumour removal. The surgical approach can be tailored to individual cases. Decompression of the optic canal can be achieved either through a combined frontotemporal approach and OZ or a frontotemporal approach alone. However, the frontotemporal approach combined with OZ in these series was not associated with better visual or proptosis outcome.

When the major symptom is proptosis without optic canal stenosis, extended lateral orbitotomy alone can be preformed to avoid the complications of a craniotomy. As the majority of SOMs are resected subtotally, postoperative radiotherapy may provide better control of tumour growth. As retrospective studies, by their nature, have potential bias, these case series should be followed in future by prospective studies.
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Table 1. Visual outcome after treatment of sphenoorbital meningiomas

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<th>Treatment Modality</th>
<th>No. of patients ((n = 66))</th>
<th>Outcome of treatment (no. of cases)</th>
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<th>Stable/No significant change</th>
<th>Significant deterioration</th>
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<td></td>
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<td>0</td>
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<td>1</td>
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<td>12</td>
<td>10</td>
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<tr>
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<td>10</td>
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</table>

| Radiotherapy       | 15                           | 0                                  | 15          | 0                          |

VA = Visual acuity; TX = Treatment; OZ = Orbitozygomatic resection
Fig. 1

A–B
Preoperative and postoperative T1-weighted MR fat suppression sequences obtained post-Gd administration, showing a sphenoorbital meningioma before and after frontotemporal craniotomy combined with orbitozygomatic resection.

C–D
Preoperative and postoperative CT scan axial views showing a sphenoorbital meningioma before and after extended lateral orbitotomy.
Figure -2 Proptosis reduction in three different surgical approaches

1. Frontotemporal craniotomy   2. Frontotemporal craniotomy combined with orbitozygomatic resection   3. Extended lateral orbitotomy