Surgical management of tuberculum sellae meningiomas: involvement of the optic canal and visual outcome

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METHODS

Objective: To present a large series of surgically treated tuberculum sellae meningiomas with particular regard to involvement of the optic canal and visual outcome.

Methods: A retrospective analysis was done on 53 patients (40 female) with meningiomas originating from the tuberculum sellae who underwent surgery between 1991 and 2002. The standard surgical approach consisted of pterional craniotomy. Sixteen meningiomas extended posteriorly onto the diaphragma sella, 29 anteriorly to the planum sphenoidale, and 19 to the anterior clino Pl. Thirty seven tumours involved the optic canal, three bilaterally. Follow up ranged from 6 to 108 months (mean 29.9 months).

Results: Total macroscopic resection was achieved in 48 patients. Median tumour size was 2.6 cm. Postoperatively, visual acuity improved in 20 patients and deteriorated in seven. Preoperative and postoperative visual acuity worsened with increasing duration of preoperative symptoms and with increasing age. Extension into the intracanal space was a negative predictor. However, tumour size did not influence visual acuity. Recurrence occurred in two cases (21 and 69 months postoperatively). Two patients died from causes unrelated to the tumour.

Conclusions: In the majority of patients with tuberculum sellae meningiomas, total resection may be achieved through a pterional approach with minimal complications.

Meningiomas of the anterior skull base account for 40% of all intracranial meningiomas. Of these, about 25% are tuberculum sellae tumours. Women are affected three times more often than men. The disease is diagnosed in the fourth or fifth decade. The tumour originates from the tuberculum sellae, chiasmatic sulcus and limbus sphenoidale, and the diaphragma sella. Tuberculum sellae meningiomas very commonly extend into both optic canals, a problem that is underemphasised in published reports. Visual loss in one eye with optic atrophy is the initial and most common symptom. These tumours tend to displace the optic chiasm backwards and the optic nerves laterally and superiorly. Compression of the chiasma is typically accompanied by bitemporal hemianopia. Different techniques for resection include the pterional approach or a unilateral or bilateral subfrontal approach. Indispensable surrounding structures such as the optic nerves and chiasm, pituitary stalk, and anterior choroidal artery must be respected.

This report summarises a retrospective analysis of 53 tuberculum sellae meningiomas treated at the neurosurgical clinic of Duisburg over an 11 year period. The study places special emphasis on surgical technique and visual outcome. Postoperative visual outcome is analysed and related to preoperative visual loss, tumour volume and topography, and to the age of the patients. Prognostic variables for vision are defined, and compression patterns of the optic nerve and chiasm elaborated. The frequency and extension of optic canal involvement and its possible negative effect on prognosis is investigated and the importance of optic canal inspection for possible tumour extension discussed.

Patient characteristics

The series consisted of 53 patients with tuberculum sellae meningiomas, all of whom underwent surgery from 1991 to 2002. The cohort comprised 40 women and 13 men (female to male ratio, 3:1). Follow up data were available from 6 to 108 months, with a median of 29.9 months. The mean (SD) age of the patients was 52.6 (13.6) years (range 27 to 78).

Ophthalmological examination consisted of testing the patients’ visual acuity (Snellen notation), fundoscopy, and Goldmann perimetry for visual field defects. A change of ≥2 lines indicated improvement or deterioration. Oculomotor function was evaluated, along with other neurological functions. Endocrinological tests (adrenal, thyroid, and gonadal axes, specific gravity of the urine, fluid balance) were done preoperatively and one week and three months postoperatively.

All patients underwent evaluation by computed tomography (CT) or magnetic resonance imaging (MRI) with and without contrast agent. High resolution CT showed hyperostosis of the planum sphenoidale and tuberculum sellae or calcification within the tumour. Both T1 and T2 weighted MRI were done in three planes to analyse the relation to vascular and neighbouring structures. Follow up MRI was scheduled at three and 12 months postoperatively, and then annually.

Information on clinical history, signs, surgical approach, and outcome was obtained retrospectively by review of the patient’s case notes and the radiological reports. All 53 meningiomas originated from the tuberculum sellae. Sixteen extended posteriorly into the diaphragmal region, 29 anteriorly to the planum sphenoidale, and 19 to the anterior clino Pl. Thirty seven involved the optic canal (fig 1). Nine extended along the optic canal into the intraconal orbital compartment. One presented as a large bulbiform orbital mass; this patient presented with one large mass at the tuberculum sellae and the second mass around the optic nerve. It is probable that this intraorbital tumour had extended into the intracranial space, forming a second tumour nodule. The size of the tumour varied from 1 cm to 5 cm, with a median diameter of 2.6 cm.
Surgical technique

All operations were carried out through a unilateral frontotemporal approach, on the side of visual deterioration. In bilateral involvement the right side was preferred. This pterional approach is well described.

Drainage of cerebrospinal fluid (CSF) was done by lumbar drainage and later by opening the basal cisterns. The sylvian fissure was routinely opened and the M1 segment of the middle cerebral artery exposed (fig 2, step 1). The internal carotid artery was identified (step 2) and this led to the ipsilateral optic nerve (step 3), which might be covered by tumour. The anterior tumour capsule was opened and the basal blood supply was interrupted by lifting the tumour and coagulating the feeding arteries (step 4) until the ipsilateral optic nerve appeared. The tumour was further debulked to reduce its volume until the contralateral optic nerve became visible (step 5). The tumour located medial of the ipsilateral optic nerve was then removed from the skull base (step 6). This step was followed by dissection of the arachnoidal plane from the ipsilateral optic nerve (step 7). After interruption of the basal blood supply, the tumour became soft and anaemic and thus could easily be detached from the arachnoidea of the gyrus rectus, the A1 segment of the anterior cerebral artery, or the anterior communicating artery. Finally, the tumour was removed from the A1 segment (step 8) and the chiasm (step 9).

Protection of the optic and chiasmatic blood supply is extremely important to preserve vision. In particular, the small vessels from the carotid artery to the optic nerve have to remain in their arachnoidal layer and should not be occluded. In cases of hyperostotic planum sphenoidale the basal dura was excised and the bone was drilled away. The defect was covered with fat, freeze dried dura, and fibrin glue. In optic canal involvement the first 3–5 mm of the optic canal were drilled away if there was further tumour extension. The ophthalmic artery below the optic nerve was exposed and preserved. If necessary, the optic nerve sheath was opened until the annulus of Zinn was reached and the tumour around the optic nerve was carefully removed. In contrast to optic nerve sheath meningiomas, tumour mass in the optic canal in tuberculum sellae meningiomas could easily be dissected from the optic nerve.

All operations were done by the senior author, WH. The technique has only slightly changed over the years. The most important steps remain the interruption of the basal blood supply and hollowing of the tumour. The most important structure to be exposed is the contralateral optic nerve to obtain an anatomical overview. Close to the optic nerves, no active coagulation was used and the tumour was sucked within its arachnoidal plane. In recurrent tumour growth no radical resection was done. Deroofing of the optic canal was carried out whenever intracanalicular tumour growth was suspected. The tumour is always located in the basal part of the canal. With deroofing of the canal, the optic nerve can be mobilised and then the tumour below the nerve can be sucked away. This procedure prevents recurrent tumour growth.

Statistical analysis

Spearman’s correlation coefficients ($r$) were calculated. Non-parametric testing was used because abnormal distributions were suggested by Kolmogorov–Smirnov normality tests. The non-parametric Mann–Whitney U test was applied to independent samples. For two dependent samples we used the Wilcoxon test, and for more than two independent samples the Kruskal–Wallis H test. For comparisons between nominal characteristics we used $\chi^2$ tests. Probability values below 0.05 indicated significant differences.

RESULTS

Signs and symptoms

The mean duration of symptoms was 21.8 months (range 1 to 240). The preoperative duration of symptoms correlated negatively with preoperative visual acuity (Spearman’s correlation coefficient $r = -0.34$, $p = 0.006$). Postoperative recovery of visual deficits was worse within the group of patients with deficits lasting longer than six months (Kruskal–Wallis test, $p = 0.005$). There was a clear correlation between age and preoperative visual disturbances (Spearman’s correlation coefficient $r = -0.42$, $p = 0.0007$). Visual impairment was the initial symptom in 40 patients,
visual field defects in four, and headache in three. Six meningiomas were an incidental finding.

On admission, seven patients had normal vision, 17 still had good vision (≥0.5), 13 had fair vision (<0.5 to >0.1), and 15 had no useful vision (≤0.1 to 0) (table 1). Location in the intraconal compartment was associated with worse preoperative vision (Wilcoxon test, p = 0.018). Other extensions from the tuberculum sellae did not influence visual acuity (Wilcoxon test, NS). Visual impairment of the other eye was present in 18 patients, among whom three had no useful vision. The tumour size correlated with contralateral visual impairment (Spearman rank correlation coefficient \( r = -0.32, p = 0.008 \)).

Fundoscopy revealed optic disk pallor in 26 patients, of whom seven showed bilateral involvement. Disc swelling was found in four patients of whom two showed bilateral swelling. All but three patients with impaired vision also had visual field defects (n = 43). Prechiasmal and anterior

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**Table 1** Overview of preoperative visual acuity (Snellen notation), visual outcome, and follow up

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Figure 2  (A) Schematic drawing of the target areas visible through a pterional trepanation (intradural structures left sided and bony structures right sided). (B) Anatomical sketch of the tuberculum sellae meningioma with adjacent structures (the brain stem and its vessels are retracted). Surgical steps: (1) identification of the M1 segment; (2) internal carotid artery bifurcation; (3) ipsilateral optic nerve and optic canal; (4) interruption of the blood supply and debulking; (5) contralateral optic nerve; (6) dissection from the skull base along the interoptic fold; (7) dissection of the arachnoidal plane from the ipsilateral optic nerve; (8) dissection from the A1 segment and (9) from the chiasm.
chiasmal compression resulted in central scotoma and temporal anopia in one eye and incomplete temporal anopia in the other eye (type 1, n = 4). Compression of the chiasm could provoke the classical bitemporal hemianopia (type 2) with reduction in central visual acuity in one or both eyes (n = 17, eight complete, nine incomplete). Involvement of the intracranial optic nerve at the junction to the chiasm led to the anterior junction syndrome (type 3) with ipsilateral central scotomas and contralateral temporal anopia (n = 5). Lesions of the optic tract resulted in incongruent homonymous hemianopia (type 4) in three patients. Three patients had irregular small unclassified visual field defects and 11 were blind in one eye. Hyposmia was present in four patients.

Tumour related endocrinological abnormalities were seen in four patients (one had a raised prolactin level, one had polydipsia, one had partial insufficiency of the adrenal axis, and one had a raised cortisol level). The patient with central Cushing’s syndrome was first operated on for the tuberculum sellae meningioma and three months later an ACTH producing intrasellar microadenoma was removed through a trans-sphenoidal approach (fig 3).

**General history**

Three patients suffered from multiple intracranial or spinal meningiomas, and two from optic nerve sheath meningiomas. Three patients had undergone operation for tuberculum sellae meningiomas in other neurosurgical departments (2, 7, and 24 years before) and were referred to our centre for reoperation. One patient was misdiagnosed as a pituitary adenoma and first operated on by a trans-sphenoidal approach.

**Surgery**

The standard approach was the intradural pterional approach. Thirty six patients were operated on through a right sided approach and 17 through a left sided approach. The arachnoid remained intact in 47 cases. Tumour size influenced the preservation of the arachnoid (U test, p = 0.017). With respect to preoperative and postoperative visual acuity, there was no difference between the groups with preserved and non-preserved arachnoid (U test, NS). However, there was a tendency to worse postoperative visual acuity in the group with non-preserved arachnoid medial to the optic nerve. The pituitary stalk could be preserved in all cases. Total macroscopic resection was achieved in 48 patients. Five patients had subtotal resection owing to small tumour rests around the vessels or the optic nerve or to bone involvement. The extent of resection did not depend on tumour size (U test, NS). The ipsilateral optic canal was explored in 37 patients. The dural part was involved in 21 cases. The bony roof was drilled away in 16 patients who had further tumour extension. Bilateral involvement of the optic canal was seen in three patients. However, the extent to which the optic canal was opened did not influence postoperative visual acuity (Kruskal–Wallis test, NS). Only deroofing of the optic canal allows total tumour removal below the optic nerve within the optic canal.

Neither the extent of resection (U test, NS) nor the size of the tumour (Spearman rank correlation, NS) influenced postoperative visual acuity.

**Complications**

Two patients (aged 77 and 78 years) died one month postoperatively from cardioembolic brain stem infarction and septicemia (overall mortality 3.7%). One patient suffered a cardioembolic left occipital infarct. One epidural haematoma required reoperation and two subdural hygromas.
were punctured. One patient had to undergo a second operation for rhinoliquorrhea from the frontal sinus. Five patients had a temporary cerebrospinal fluid fistula and were adequately treated by lumbar drainage. Four suffered from an epileptic seizure in the first postoperative week. Secondary visual loss in the right eye occurred in one patient on the 12th postoperative day; he underwent treatment with nimodipine and haemodilution therapy, with complete recovery within five days. One patient received hydrocortisone as replacement therapy for three months. Four patients developed permanent hyposmia and one developed anosmia.

Outcome
On discharge, the absolute postoperative visual acuity improved in 20 patients (≥2 lines in Snellen notation), remained unchanged in 25, and deteriorated in seven. Preoperative vision correlated highly with postoperative vision (Spearman’s correlation coefficient \( r = 0.69, p = 4.9 \times 10^{-9} \)). Age was negatively correlated with the postoperative visual acuity (Spearman’s correlation coefficient \( r = -0.34, p = 0.006 \)). Longer duration of preoperative symptoms correlated with worse recovery (Spearman’s correlation coefficient \( r = -0.47, p = 1.85 \times 10^{-7} \)). Deterioration of vision was not associated with involvement of the optic canal. In these cases, there was a tendency to interruption of the arachnoidal plane. Thus deterioration of visual acuity seems to be a vascular problem.

Eighteen patients with visual field defects improved markedly postoperatively. In eight patients the defects resolved completely. However, seven patients worsened. No patient developed diabetes insipidus.

Follow up
Vision generally improved rapidly within the first weeks after surgery, but continued to improve for the next year. At follow up, the absolute visual acuity improved further in four patients and worsened in four. Only one patient with bad vision (<0.1 to 0) improved further. Patients older than 50 years had worse recovery (U test, \( p = 0.005 \)). Vision did not deteriorate with longer follow up (Spearman’s correlation coefficient, NS).

Recurrence
Recurrence was found in two cases (3.7%), 21 and 69 months postoperatively. However, follow up varied from 6 to 108 months. One of these patients became symptomatic with a new visual field defect and the other with headache. The case with early recurrence had a history of multiple meningiomas and the 4 cm tumour was only subtotally resected at the first operation. Both patients underwent reoperation with good results.

DISCUSSION
Outcome and prognostic factors
The major visual signs associated with tuberculum sellae meningiomas are visual loss (98%) and optic atrophy (78%). The Foster–Kennedy syndrome of unilateral optic atrophy and contralateral papilloedema occurs in 5% of these patients.1 The clinical presentation is of visual loss in one eye with optic atrophy that develops into blindness and diminution of acuity in the opposite temporal visual field.1 About 80% of patients suffer from visual disturbance for a period of between three months and 20 years, with an average of more than two years.4

Hemiparesis is seen in 15% and anosmia in 11%. Pituitary insufficiency has been reported to occur in up to 22% of patients.1 However, in the study by Fahlbusch and Schott5 endocrine disturbances did not play a major role, but may be a more important factor in very large tumours.

Improvement of vision was noted in 50–66%, while in 17–28% it remained unchanged. Deterioration was noted in 10–25%.6–11 Tumours larger than 3 cm2 or 4 cm3 are thought to be associated with increased morbidity and a lack of improvement in vision following surgery, as well as with increased mortality.1 Visual outcome was favourably affected when the tumour was less than 3 cm, the preoperative visual loss was less than 50% and present for less than two years, and the optic discs appeared normal on fundoscopy.11 Poor prognostic factors include tumour extension into the optic nerve or the diaphragma sellae. Fahlbusch and Schott reported a markedly increased risk of visual deterioration (up to 40%) in patients in whom the duration of symptoms was longer than one year.

In the series by Zeygaridis et al.,12 visual prognosis was favourably affected by age under 54 years, duration of symptoms of less than seven months, and the presence of an intact arachnoid membrane around the lesion. Severe preoperative loss of visual acuity appeared to be an unfavourable prognostic factor. Severe morbidity occurred in 6.4%. However, in our study tumour size and location had no effect on outcome, in contrast to some previous reports. Our findings are supported by recent studies.13

Visual recovery is sometimes possible in patients with poor preoperative visual acuity (eight of our patients in this group improved, and six even regained good vision). Postoperative severe deterioration is mostly permanent. Only one patient with secondary visual loss in one eye on the 12th postoperative day recovered completely.

The incidence of total removal of tuberculum sellae meningiomas has varied greatly, with reported rates of 40–95%.1 Mortality ranges from none to 7%.1 A large study on recurrence of cranial base meningiomas13 showed a perioperative mortality of 10.8%, and 9.7% died within 10 years. Our mortality was 3.7% in our oldest patients. Cornu et al.14 found that the general condition of the patient was more important than age itself. Patients in poor general condition had a fivefold increased risk of a poor outcome.

Optic canal involvement
Bilateral involvement of the optic canal is described.1 Growth of tumour in the optic canal is thought to be associated with worse visual prognosis.15 Arai et al.15 emphasised that the presence of preoperative visual disturbance implies the need for optic canal deroofing. Two of 13 cases required deroofing on both sides because the tumour extended into the canal on the non-symptom side. Four of eight cases with bilateral visual disturbance required optic canal deroofing on both sides. Andrews and Wilson16 noted that patients with invasion of the optic canal show less frequent improvement in visual acuity or visual fields. DeMonte1 favours optic canal inspection at the end of the surgery. Residual tumour in the canal may be the site of recurrence or the cause of a lack of visual improvement following surgery.1

In our opinion, when the tumour is in the optic canal it should be followed and removed completely, together with any dura involved. Tumour within the optic canal can easily be removed because there are no adhesions. However, it may be difficult to dissect the tumour from the chiasm and nerves within the cistern. An intracanal tumour location had a negative effect on visual outcome. The high proportion of intracanalicular extension (69.8%) supports our procedure of surgical inspection and tumour removal.

Pathogenesis
There are different mechanisms for preoperative optic nerve injury: ischaemia, compression, demyelination, and tumour invasion.17 Compressive mechanical injury leads to small vessel compromise and demyelination, especially in patients...
with a long duration of visual loss before surgery. Assuming that there is no additional intraoperative trauma to the optic nerve, incomplete or no recovery of visual function after surgery may imply chronic severe preoperative ischaemic or compressive damage and demyelination. Compressive injury can at least be reversed by surgery. Thus decompression of the optic pathway is the main goal of the surgery. Zeygiridis et al. hypothesise that ischaemia is the cause of primary visual loss rather than simply optic nerve compression. Younger patients are able to compensate for microvascular deprivation more effectively than older ones. This could explain our finding of worse vision and poor recovery in elderly patients. Andrews and Wilson postulated direct compression or invasion of the nerve, or preoperative damage to its blood supply. When the tumour is densely adherent to the optic apparatus, the best choice is to leave tumour fragments. Preservation of the vascular supply to the nerves is important. Occlusion of perforating vessels contributes to postoperative visual deterioration and should be avoided.

**Surgical approach**

Symon and Rosenstein reached the tumour subfrontally along the midline following right fronto craniotomy. Al-Mefty et al. chose unilateral or bilateral frontal craniotomy, depending on the tumour size. Ohta et al. used the pterional approach in 15 patients, the orbitozygomatic in 10, and the bilateral subfrontal approach in six. Arai et al. describe a transcranial trans-sphenoidal approach for tuberculum sellae meningiomas. They operated on 21 patients through a bifrontal craniotomy, removal of the orbital rim, transsection meningiomas. They operated on 21 patients through a transcranial trans-sphenoidal approach for tuberculum sellae meningiomas. Only in the series of Fahlbusch and Schott was radical surgery, and 25–45% for patients undergoing subtotal removal. The five year recurrence rate was 4% for patients undergoing subtotal removal. The recurrence in subtotal removal occurred earlier. Complete tumour resection should not be achieved at the cost of increased morbidity. On the other hand, surgical resection should be as complete as possible to avoid the risk of future recurrence.

**Radiotherapy**

Becker et al. reported a series of 24 secondary optic nerve sheath meningiomas undergoing stereotactic fractionated radiotherapy (SFRT) using 6 MV photons from linear accelerators. The median total tumour dose was 54 Gy using 1.8 Gy/fraction. The visual fields were improved in six of 24 examined eyes and visual acuity in seven of 26 in this series. Stable visual fields and stable visual acuity were observed in 17 of 24 and 19 of 26 patients, respectively, without toxicity. Secondary optic nerve sheath meningiomas were defined as arising from the intracranial meninges and subsequently involving the visual pathways. This means that they include all kinds of skull base pathologies such as medial sphenoid wing meningiomas, cavernous sinus meningiomas, planum sphenoidale meningiomas, and perhaps also some tuberculum sellae meningiomas.

Other large series of cases receiving radiotherapy for skull base meningiomas have emphasised the stabilisation of the tumour volume with good clinical results. However, only one series—that of Pend—incorporated three sellar meningiomas. The high efficacy of SFRT for large skull base meningiomas and the long term results have also been reported by Debus et al. Their overall survival rate was 97% after five years and 96% after 10 years (WHO grade I). Local treatment failure was observed in three of 180 patients with WHO grade I tumours and in two of nine patients with grade II tumours. A volume reduction of >50% was observed in 14%. Pre-existing cranial nerve symptoms resolved completely in 28% and treatment induced toxicity occurred in 1.6%.

We think that radiotherapy should be considered in elderly patients and in those with subtotally resected tumours. Nevertheless, surgery remains the treatment of choice in tuberculum sellae meningiomas.

**Conclusions**

The goals of surgery are tumour control, minimal morbidity, and visual improvement. The duration of preoperative symptoms has a strong influence on visual outcome. Thus early diagnosis is desirable for successful treatment with a better chance of good postoperative visual outcome. Tumour extension into the optic canal is quite common and required optic canal deroofing and gentle tumour removal.

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**REFERENCES**

Bilateral first rib fractures due to tardive dystonia

A 37 year old man with mild learning disability and a history of previous neuroleptic use was admitted for investigation of a 1 year history of dystonia. On admission he had severe axial and limb dystonia with opisthotonus. Following extensive investigation for other causes of dystonia, a diagnosis of tardive dystonia secondary to neuroleptic exposure was made.

A routine chest radiograph revealed bilateral first rib fractures (fig 1). Such fractures are extremely rare because the first ribs are deeply placed and well protected by the shoulder girdle, lower neck musculature, and clavicles, and almost always occur in the context of severe, generalized trauma. However, rare cases of non-traumatic first rib fractures caused by sudden violent contraction of the neck muscles or repeated muscular pulling, for example during fractures caused by sudden violent contraction of the neck trauma. However, rare cases of non-traumatic first rib fractures rarely occur in the context of severe, generalized

The patient reported here had no history of previous trauma, and had no abnormal physical signs in the chest, nor any distal neurovascular deficit in the arms. We propose that the bilateral first rib fractures in this case resulted from tension on the scalene muscles during repetitive, violent opisthotonic spasms. To our knowledge, this is a unique neurological cause of bilateral first rib fractures.

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Competing interests: none declared

NEUROLOGICAL PICTURE

Bilateral first rib fractures due to tardive dystonia

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References

Figure 1 Chest radiograph showing bilateral first rib fractures (arrowed).

Surgical management of tuberculum sellae meningiomas: involvement of the optic canal and visual outcome

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