Case Report

Delayed Visual Loss Following Transcranial Surgery for Tuberculum Sella Meningioma

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INTRODUCTION

Postoperative visual loss is a dreaded complication following transcranial excision of tuberculum sella meningioma. Visual deterioration is commonly noticed immediately after surgery and delayed deterioration after 72 hours is uncommon. We report a case of delayed postoperative deterioration in a 48-year-old lady and discuss the possible mechanisms.

CASE REPORT

A 48-year-old female presented with 1 year history of headache and progressive blurring of vision of the right eye. On examination, she had only perception of light in her right eye, and fundoscopy revealed primary optic nerve atrophy in the right eye. The left eye vision and fundus examination were normal. Imaging [Figure 1] revealed a suprasellar enhancing dural-based lesion suggestive of a tuberculum sella (TS) meningioma. She underwent right pterional craniotomy and Simpson’s grade II excision of the meningioma along with intradural optic canal deroofing. Postoperative recovery was uneventful, and she continued to perceive light in her right eye. On the 7th day, she complained of complete blindness of her right eye. Repeat imaging was not contributory. Fundoscopy showed a pale atrophic disc. She was treated with steroids for 5 days. Her vision failed to improve, and 6 months after surgery, she remains blind in her right eye.

DISCUSSION

Suprasellar meningiomas constitute 5%–10% of all intracranial meningiomas. TS meningioma refers to tumors arising from tuberculum, limbus sphenoidale, chiasmatic sulcus, and diaphragm sella. TS meningiomas generally present with visual symptoms, and deterioration of vision is the most common indication for surgery in patients with TS meningioma. Analyzing the anatomical origin of their 100 cases of TS meningiomas, Kim et al. have proven that these tumors have their point of origin laterally near one of the optic canals in >85% of cases and then proceed toward the midline. Visual symptoms can thus be asymmetric and could be either from mechanical compression from the tumors itself or impingement of the nerve in a tight optic canal. Considerable debate exists regarding the ideal mode of approach and the extended endoscopic transsphenoidal approach seems to be increasingly becoming popular. Comparing endoscopic endonasal versus open transcranial resection of TS meningiomas, Komotar et al. observed that the visual outcome is better with an inferior approach – both in terms of improvement of existing deficits as well as in terms of postoperative deterioration.[1] In terms of extent of resection and risk of cerebrospinal fluid leak, the transcranial approach has an edge and still remains the most common surgical approach for TS meningiomas.[1]

Different studies have shown a 58.4% of improvement and 12.9% risk of postoperative deterioration.

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following transcranial surgery. Visual deterioration is commonly noticed immediately after surgery and is linked to multiple reasons. These include intraoperative manipulation of the nerve, thermal injury due to use of cautery and drills and any obvious vascular injury. In the authors’ personal series of 71 TS meningiomas, postoperative worsening of vision occurred in 3 (4.22%) cases and was apparent immediately postoperatively in all the cases. However, in this case, the visual deterioration was noticed at the time of discharge 7 days after surgery.

The reasons for delayed deterioration remains unclear in our case. In a similar report of two patients, Santarius et al. postulate an ischemic etiology due to vasospasm. They performed angiography and verapamil infusion in their patients, and although angiography did not reveal any significant vasospasm, verapamil infusion did help in recovery in one of the patients. Santarius et al. have thus evolved a protocol of hypervolemic hypertensive therapy for 48 h postoperatively for all tuberculum and periorbital meningioma patients in whom the clinoid or optic canal is drilled. Their postulate being that during surgery a critical quantum of optic pathway-splying vasculature is compromised making the tissues vulnerable to ischemia, and it would be ideal to avoid further tissue hypoperfusion by ensuring adequate blood pressure.

Venous insults manifest in a subacute fashion unlike an arterial ischemia, and occlusion of venous drainage could be related to delayed deterioration. Preservation of veins in the suprasellar area is not carried out as diligently as in other areas such as parasagittal area. Our case did have a small venous injury which was thought to be inconsequential at the time of surgery. Venous drainage of the optic apparatus probably needs to be studied more, and all attempts should be made to preserve them whenever possible.

It is well known that decompression of a critical spinal canal stenosis can result in postoperative neurological deterioration secondary to spinal cord edema. Probably, a similar mechanism with sudden decompression resulting in probable reperfusion injury could happen after optic nerve decompression. We believe that more the severity of compression, more the chances of deterioration with decompression. Thus, patients with significant preoperative optic nerve compression as evidenced by severe optic atrophy and minimal vision (hand movement or perception of light) are more likely to have postoperative deterioration as was seen in our patient.

Thermal injury due to cautery and drills and mechanical injury due to manipulation could at times result in delayed optic nerve edema which could be another possible etiology for delayed visual deterioration; most of the TS meningiomas have an optic canal extension, and Mortazavi et al. have suggested early decompression of the optic canal by extradural clinoidectomy and optic canal deroofing to reduce intraoperative manipulation of the optic nerve. This along with careful use of cautery with liberal irrigation of saline while coagulating the dural attachment are some maneuvers which can prevent postoperative visual deterioration.

**CONCLUSION**

Postoperative visual deterioration is not an uncommon sequel following transcranial excision of a TS meningioma. The deterioration can occur immediately or few days postoperatively. Patients with severe optic atrophy and poor vision are more likely to have this complication. The possible mechanism could be vascular, mechanical, or sequel to sudden decompression. Extradural clinoid drilling, adequate hydration during the use of drill and cautery, gentle and careful handling of optic nerves, intraoperative use of papaverine, and postoperative hypervolemic hypertensive therapy are a few prophylactic steps which could prevent this catastrophe.

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There are no conflicts of interest.

**REFERENCES**

