Planum Sphenoidale Meningioma: A Rare Location of Skull Base Meningioma

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Abstract

Background: Meningioma is common primary central nervous system tumors. Twenty-five percent of all meningioma consist of skull base meningioma. Planum sphenoidale meningiomas are rare. Planum sphenoidale meningiomas can extend into adjacent areas. Approximately two thirds of patients complain of failing vision in one eye as the first symptom.

Case Report: A 32-year-old woman presented with 6-month history of progressively worsening blurred of both vision. She also complained her smell ability was reduced for 3 months. She had headache for 6 months. The pain was worsening in the morning. She is conscious. A neurologic examination revealed bilateral hyposmia and visual deficits but no weakness. Visus of oculo dextra was 1/300 and visus of oculo sinistra was no light perception. Magnetic resonance imaging (MRI) intravena contrast of brain revealed a large extra-axial mass measured ±6,2x5,9x6 cm centred on planum sphenoidale displacing both frontal lobes. She had an operation of tumor removal with cranio-orbitozygotomy approach. The tumor, which measured ±7cmx7cmx6 cm, was successfully removed completely. She gets improvement of smell ability and both visual postoperatively.

Discussion: Planum sphenoidale meningiomas present a frequently encountered pathology of the anterior skull base. These meningiomas give rise to an early visual disturbance with relatively slow progression. Displacement of the olfactory tracts and optic chiasm occur when the meningioma extends into the paranasal sinuses and nasal cavity. Clinical presentation and diagnosis often occur in the late stage. Anosmia is one of common finding on physical examination. Postoperative improvement of visual symptoms depends on the preoperative duration of those symptoms.

Conclusion: Planum sphenoidale meningiomas are rare cases. Headache, visual deficits and impaired smell are common clinical presenting.

Keyword: Planum Sphenoidale Meningioma, Visual deficits, Anosmia, Neurooncology

Introduction

Meningioma account for approximately 36% of all primary central nervous system tumors, representing the most frequently diagnosed primary brain tumor. Twenty-five percent of all meningioma consist of skull base meningioma.[1] Planum sphenoidale meningiomas are rare, constituting only 2% of all primary intracranial tumours.[2] They are benign, slow-growing tumours.[3] Planum sphenoidale meningiomas can extend into adjacent areas such as the diaphragm sellae, posterior clinoid and superior wall of the cavernous sinus. Complete meningioma resection including vascular supply and involved dura and bone may be difficult due to their intimate involvement with these nearby structures.[4] In most patients visual loss has an insidious onset, and the course is progressive. It may, however, be acute or fluctuating.

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Approximately two thirds of patients complain of failing vision in one eye as the first symptom, and monocular blindness may be present in half of the patients before surgery.[5]

**Case Report**

A 32-year-old woman presented with 6-month history of progressively worsening blurred of both of vision. First it affects right vision then followed left vision. Left vision is loss. She also complained her smell ability was reduced for 3 months. She had headache for 6 months, which reduced by analgetic. The pain was worsening in the morning. She consequently lost his job as her poor performance. She needed someone to take care of herself. She had no behavioural changes and no past medical history of illness or known exposure to ionizing radiation. She had no known family history of intracranial tumor.

On examination, she is conscious. A neurologic examination revealed bilateral hyposmia with visual deficits and no weakness. Visus of oculo dextra was 1/300 and visus of oculo sinistra was no light perception. Initial blood investigation results (ie complete blood count, glucose level, liver function, renal function, thyroid function test) were normal.

Magnetic resonance imaging (MRI) intravena contrast of brain revealed a large extra axial mass measured ±6,2x5,9x6 cm centred on planum sphenoidale displacing both frontal lobes (Figure 1)

![Figure 1. MRI Brain IV Contrast](image)

Mass effect and entrapment of the left lateral ventricle were also visible. She had an operation of tumor removal with cranio-orbito-zygotomy approach (Figure 2). The operation took about 5 hours long. The tumor, which measured ± 7cm x 7cm x 6cm, was succesfully removed completely (Figure 3). She was conscious postoperatively.
Figure 2. Exposed tumor   Figure 3. Macroscopic of tumor

Post operatively, the ability of smell was increasing then before even. She had presented both visual improvement that visus of both oculi become 1/60. After 10 days, she went home and followed in outward. The histopathology of the tumor revealed meningioma WHO grade I (Figure 4).

Discussion

Planum sphenoidale present a frequently encountered pathology of the anterior skull base, representing according to some authors 5 to 10% of all intracranial meningiomas.[5] These meningiomas give rise to an early visual pathology with relatively slow progression, but due to the fact that other symptoms are missing or are subtle they have a larger tendency to develop undiagnosed for longer periods of time.[6]

Displacement of the olfactory tracts and optic chiasm occur when the meningioma extends into the paranasal sinuses and nasal cavity.[3] Clinical presentation and diagnosis often occur in the late stage, as many patients are asymptomatic before the meningioma reaches a sufficient size (> 4 cm) to compress the frontal lobe and optic nerve or optic
chiasm.[7] The patient usually presents with dysexecutive syndrome (severe cognitive impairment and profound changes in personality), which is usually first noticed by family members; headache; or visual symptoms. Anosmia is a common finding on physical examination, but it is not a typical presenting symptom.[8] Postoperative improvement of visual symptoms depends on the preoperative duration of those symptoms.[9] Loss of olfactory function is usually permanent.[3] Although MRI is more expensive and less readily available, it is preferred to CT scan.[10]

Similarly, once tumors approach <6 mm from the optic chiasm, they are 11 times more likely to cause visual symptoms, implying at first that it might be advisable to operate on a planum meningioma that is 7 to 8 mm away from optic chiasm even if it is not currently causing visual symptoms. Interestingly, however, distance to the optic chiasm is not an independent predictor of visual symptoms in our multivariate analysis. Only brain tumor volume is an independent predictor of visual symptoms in our multivariate model. Although this may result from our relatively limited sample size and we certainly believe it is advisable to consider the proximity to the chiasm in treatment decision planning, our data raise the possibility that it not contact with the optic nerve alone that causes visual symptoms but rather overall tumor bulk. Patients with large tumors may have chronic papilledema from increased intracranial pressure leading to nerve loss and optic atrophy on funduscopic examination.[11]

Some patients may present first with visual loss and acute hemorrhagic papilledema leading to rapid visual decline from ischemic optic to rapid visual decline from ischemic optic neuropathy. Some of these patients will have continued visual decline after removal of their tumors from the irreversible ischemic process at the optic nerve head that surgical decompression cannot immediately reverse, which suggests that it is important to follow visual function and field in patients with preoperative visual loss, which was not done systematically in our series because of referral patterns.[11]

Conclusion

Planum sphenoidale meningiomas are rare location of skull base meningioma. Headache, visual deficits and impaired smell are common clinical presenting. Postoperative improvement depends on preoperative duration of symptoms.
References


