A rare type of tumor: orbital schwannoma. Case report and literature review

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Abstract: Schwannomas arise in the cells responsible for the myelinating the neurons distal to the Obersteiner-Redlich zone. Most of the intracranial Schwannomas are in the posterior fossa, developed from the VIIIth or Vth nerve sheath. The location on other cranial nerves is quite rare, only 6% of the orbital tumors being Schwannomas. We review the case of a 52 years old male patient, presenting for right eye exophthalmia and visual field deficit, diplopia due to VIth nerve paresis, and stubbing pain in the right eye, the MRI showing a tumor, located in the orbital apex displaced the globe forward and superiorly, and the optic nerve medially and superiorly. A modified lateral orbital approach was preferred. The choice of the orbitotomy allowed us to maintain the integrity of the lateral rim of the orbit without the need of a bony reconstruction at the end of the intervention, as it faced the anterior margin of the temporal muscle, covering it and not the skin over the zygomatic bone. The technical approach for orbital schwannomas should be tailored to reach the lesion through an esthetic incision and orbitotomy, immediately under the resected bone, with no need retracting the ocular globe or the vasculonervous elements in the orbit.

Key words: Schwannoma; lateral orbital approach

Introduction

Schwannomas arise in the cells responsible for the myelinating the neurons distal to the Obersteiner-Redlich zone. Most of the intracranial Schwannomas are in the posterior fossa, developed from the VIIIth or Vth nerve sheath. The location on other cranial nerves is quite rare, only 6% of the orbital tumors being Schwannomas. These tumors become symptomatic through the mass effect and usually the symptoms progress slowly over a few years mirroring the slow rate of growth.

In a large series of 308 orbital tumors published in 1984 by J. Maroon, the authors counted only 5 neurofibromas, the most often encountered orbital tumors being in order
metastasis, dermoid tumors, meningiomas and cavernomas.

Most patients with orbital tumors present with slowly progressive proptosis. A partial field cut, Blurred vision and Diplopia and papillary edema with or without optic atrophy can also be present though less frequent.

Differential diagnosis with other tumors or tumor like lesions should be obtained. The most common possible lesions resembling neurinomas are meningiomas, cavernomas and dermoid tumors, each presenting characteristic features on MRI which should be sought with attention.

The relation of the tumor with the optic nerve and the oculomotor muscles must be understood from the MRI images before proceeding to resection.

The tumor may be inside or outside the muscles cone, in the orbital apex or more excentrically situated. It may encircle the optic nerve or just displace it. The relation of the tumor with the orbit walls is very important in choosing the approach as the shortest way with the minimal risk of collateral damage should be chosen. In 1941 Dandy introduced the subfrontal approach and Nafzinger used the pterional Transcranial approach for orbital tumors.

Over time with the tendency towards minimal invasiveness the extracranial approaches became more and more favored to Transcranial approaches. These include lateral orbitotomy for tumors located lateral and basal to the optic nerve and in the orbital apex, transetmoidal for the extraconal tumors located medial to the, transmaxilary for basal lesions sitting on the maxillary sinus, or transconjunctival for basal intraconal tumors.

The transcranial approaches remain in use nowadays for tumors developed both intracranian and intraorbitar or for Tumors involving the optic canal.

Case report

We review the case of a 52 years old male patient, presenting for right eye exophthalmia and visual field deficit, diplopia due to VI\textsuperscript{th} nerve paresis, and stubbing pain in the right eye. These symptoms evolved progressively during a few years. No signs and symptoms of Recklinghausen disease were recognized. MRI scan performed on admission showed a large inhomogeneous orbital tumor with central round areas of hypo intensity in both T1 and T2 and peripheral Gadolinium enhancement. The tumor, located in the orbital apex displaced the globe forward and superiorly, and the optic nerve medially and superiorly.
Figure 1 - Preop MRI showing a mix signal inferolateral orbital tumor that displaces superolaterally the optic nerve

We preferred a modified lateral orbital approach. Using an s shaped incision starting in the eyebrow and continued over the zigoma, we retracted the temporal muscle posteriorly and used a burr hole medial to the frontal process of the zigoma in the angle between the lateral and the inferior orbital walls. As this burr hole was performed over the tumor, only little enlargement of it with rongeurs was needed to mobilize the lesion circumferentially. Using a small dissector, an extracapsular resection was performed detaching the tumor from the right lateral muscle displaced superiorly and from the optic nerve, displaced superomedially. The tumor was encapsulated, of a firm consistency and a pale white color. Fortunately it was not attached to the surrounding structures and the capsule was easily dissected using a blunt instrument followed by a single piece excision of the lesion.

The choice of the orbitotomy allowed us to maintain the integrity of the lateral rim of the orbit without the need of a bony reconstruction at the end of the intervention, as it faced the anterior margin of the temporal muscle, covering it and not the skin over the zigomatic bone.

Figure 2 - Enlarging the lateral orbital wall orbitotomy with a rongeur

Figure 3 - Extracapsular dissection of the tumor with a blunt instrument

Figure 4 - Total resection in a single piece
Following an intradermal suture an ointment dressing was applied on the operated eye.

The postoperative course was uneventful, the exophthalmym resolved almost completely as demonstrated on the postop. CT. and the visual field deficit improved.

Discussion

The pathology diagnosis came much as a surprise as we expected a more frequent type of tumor like a metastasis, a meningioma or even a cavernoma yet the initial diagnosis was confirmed by the classic palisading pattern of tumor nuclei showed both on Hematoxilin-eosine and S100 stain at the final exam.

Most of the orbital Schwannomas develop from branches of the ophthalmic division of the trigeminal nerve thus being located superior and laterally or superior and medially to the optic nerve as described in an 18 years span study on 49 orbital Schwannomas. The inferolateral location of the presented Schwannoma indicates the abducens nerve as the site of origin.

As the optic nerve is mielinated in a central like fashion from oligodendrocytes and not from schwan cells, it has been hypothesized that they arrire from the sympathetic plexus inervating the vessels feeding the optic nerve. (9, 10, 13)

We described a modified Krohnline approach with incomplete resection of the frontozigomatic arcade and an orbitotomy in the angle between the lateral and the inferior wall of the orbit after a good Posterior retraction of the temporalis muscle. A classic lateral orbital approach would have been centered not on the tumor but on the superiorly displaced lateral rectus muscle. (2, 5, 6)

Most of the extracranian approaches to the orbit are based on the classical lateral orbitotomy described by Kronlein in 1889. Depending on the extension of the orbitotomy the lateral approach can open an extended though narrow surgical corridor from the lateral orbit to the superior orbital fissure, the optic canal and the cavernous sinus. (11, 12, 14) Since the use of the CT scan and MRI this tumors are diagnosed while they are confined to the orbit compared to the past when most of the orbital tumors presented an intracranian extension, henceforth the need for a transcranian approach either subfrontal preferred by Dandy or pterional preferred by Nafzinger. (3, 7, 8)

The relation of the tumor with the optic nerve is the main thing to decide the approach
as one should avoid an approach that crosses the ON in its way to the tumor. In the presented case the tumor was behind the globe, under and lateral to the optic nerve.

In 1993 a neurosurgical-ophtalmological team from Duisburg described a transconjunctival approach for an intraconal orbitobasal large cavernoma. This technique allows tumor resection without bone or muscle resection.

A transconjunctival approach could be taken into consideration in our case had the tumor been larger and bulging anteriorly from under the globe.

The superior transcranian, transorbital roof approach and the endoscopic transetmoidal approach were not considered at all as the tumor was inferior and lateral to the optic nerve. As the tumor was intraconal with no intracranial or cavernous sinus extension there was no need for a transcranial approach. (1, 4)

Another combined approach with the help of an ENT surgeon is the transmaxillary one, well suited for our case as the tumor was seated on the roof of the maxillary sinus.

Conclusion

Even if very rare, orbital Schwannomas are well encapsulated tumors, which makes the resection simple and safe. The technical approach used should be tailored to reach the lesion through an esthetic incision and orbitotomy, immediately under the resected bone, with no need retracting the ocular globe or the vasculonervous elements in the orbit.

References