Case Report

A 32-year-old lady came to our outpatient department with complaints of tightness and discomfort in the right eye, especially on looking toward extreme right side since 4 months. The tumor was totally excised with functional preservation of the nerve by superior orbitotomy. The clinical, radiological features and the management are discussed.

Key words: Abducens, intraconal, intraorbital, schwannoma

Introduction

Schwannomas are tumors arising from the schwann cells of the neural sheath of motor and sensory nerve fascicles and present as well-defined eccentrically placed masses. They may present in three clinical forms; localized schwannomas, in association with neurofibromas (as a part of von Recklinghausen syndrome), or as schwannomatosis.[1]

Intraorbital schwannomas may arise from different nerves; sometimes, it may be difficult to identify their true origin.[2] Those arising from the sensory branches of trigeminal nerve are located in superior and medial quadrants of orbit,[3] those arising from infraorbital nerve are in inferior quadrants[4] and schwannomas arising from the motor nerves usually arise at their respective myoneural junction. The literature search has identified only three other abducens nerve schwannomas of the orbit.

Case Report

A 32-year-old lady came to our outpatient department with complaints of tightness and discomfort in the right eye, especially on looking toward extreme right side since 4 months. Her visual acuity was 6/6 as per Snellen chart and visual fields symmetrical in both eyes on clinical examination. Fundoscopy revealed no abnormality. There were no neurocutaneous markers of neurofibromatosis. Patient was reassured and was asked to review after 15 days. Patient was not relieved of her symptoms, and she noticed mild proptosis of right eye. She underwent magnetic resonance imaging (MRI) scan of the orbit with contrast.

Magnetic resonance imaging scan [Figures 1-3] revealed, a 2.1 cm × 1.8 cm × 1.7 cm well-defined, lobulated intraconal mass lesion in the posterior right orbit in close proximity to the optic nerve, displacing the nerve supero-medially. The lesion is homogeneously hypointense to gray matter on T1-weighted (T1-W) and isointense on T2-W-images. On early postcontrast images, the lesion enhances homogeneously and shows intense enhancement on delayed images.

Patient underwent right frontal craniotomy and superior orbitotomy. Minimal subfrontal dural retraction was necessary. Under the operating microscope, Tenon's fascia was opened in cruciate fashion and opening extended posteriorly till the anterior margin of the optic canal. The intraconal fat was sharply dissected till the superior rectus muscle (SRM) was seen, and dissection continued medial to SRM till the superior branch of the ophthalmic artery was safely isolated. The dissection continued in this plane till the tumor was seen. The tumor was moderately vascular, soft to firm, pinkish with thick capsule. The capsule was incised, and tumor debulked...
with cavitron ultrasonic aspirator till very minimal tumor was left attached to capsule. There was no intracranal retraction during the surgery; the lateral rectus muscle (LRM) was seen with the abducens nerve innervating the muscle and tumor capsule arising from the myoneural junction. The tumor capsule was completely excised without damaging the nerve. Operative video is submitted for interested viewers [Video 1]. The edges of Tenon’s fascia were just approximated as water tight closure was not possible. The superior orbital plate was small and thin, and roof of the orbit was reconstructed with bioresorbable polycaprolactone (PCL) scaffold, which is known to have osteo-inductive properties. The scalp incision was sutured, and suction drain was placed in subgaleal space.

The histopathology [Figure 4a-d] revealed a well-circumscribed, thin encapsulated neoplasm composed of predominantly interwoven fascicles of spindle cells with elongated nuclei, eosinophilic cytoplasm, and indistinct cytoplasmic borders (Antoni A pattern). Prominent nuclear palisade is observed which is referred to as Verocay bodies. Furthermore, a focal hypocellular area with a loose, vacuolated stroma was observed (Antoni B pattern). The histological features were diagnostic of schwannoma.

Six months after surgery, patient is having complete range of extraocular movements [Figure 5] and the eyes, facial contour is symmetrical. Six months postoperative computed tomography scan [Figure 6] confirmed complete excision; bone formation was not yet visible in the PCL scaffold which was used for orbital roof reconstruction.

**Discussion**

Though abducens or sixth cranial nerve has the longest intracranial course, schwannomas of the abducens nerve are extremely rare tumors.[5] The tumor may be located within the cavernous sinus[6] or more often at the preoptine[5] region and as intrinsic brainstem tumors.[7] The literature research has identified only three other cases of isolated schwannoma of the orbit, arising from the terminal branches of the abducens nerve to the LRM.[1,8,9]

Correct diagnosis and accurate definition of the location and the extent of the lesion greatly facilitate successful surgical excision. Orbital schwannoma on MRI is usually described as a lesion that produces a low signal on T1-W images and a high signal on T2-W images, which can be homogeneously or heterogeneously enhanced.[10] The histology of these lesions underlies the variation in their appearance on MRI. Intraorbital abducens schwannoma is situated in lateral intraorbital quadrant, medial to LRM and displaces the optic nerve medially and upwards.[1,8,9]

We summarize the main features by comparing our case with those described by Irace et al.,[9] Miguel et al.[1] and Feichtinger et al.[8] [Table 1].

Similar to the description by Irace et al.,[9] Miguel et al.[1] and Feichtinger et al.[8] in our case, we were able to demonstrate preoperatively by MRI [Figures 1-3] and intraoperatively by identifying the area of attachment of the tumor, the origin of the schwannoma from the abducens nerve.

The lesions in the lateral quadrant of the orbit can be approached by lateral or superior orbitotomy. Through lateral orbitotomy Irace et al.[9] and Feichtinger et al.[8] could achieve only partial excision of solid tumor and Miguel et al.[1] could do total excision as, fortunately, the tumor was cystic. The superior orbital approach involves frontal craniotomy followed by superior orbitotomy to gain access, and it can be extended for more posterior lesions. Superior orbitotomy gives direct access to superior, medial, and lateral quadrants of orbit with minimal retraction of the intracranial structures thereby ensuring safe and complete excision of the lesions.

Based on the work of Erdogmus et al.,[11] Irace et al.[9] proposed that the entry point of nerve fibers in the muscle (e.g. area nervosa or myoneural junction)[12] is the zone where intraorbital schwannomas truly arise. When performing intraorbital surgery, anatomic knowledge of the innervation of the extraocular muscles allows one to minimize damage to neural structures. In particular, when approaching the superolateral compartment of the orbit, the knowledge of the course and branching of the abducens nerve is required. This cranial nerve runs on the medial surface of the LRM and innervates in the middle third of the muscle. In our case, this knowledge was essential to preserve nerve integrity.

### Table 1: Patients with intraorbital abducens nerve schwannoma

<table>
<thead>
<tr>
<th>Author</th>
<th>Age/sex</th>
<th>Clinical presentation</th>
<th>Tumor size (cm)</th>
<th>Tumor type</th>
<th>Surgical approach</th>
<th>Extent of resection</th>
<th>VI nerve function</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Irace et al.,[9]</td>
<td>55/male</td>
<td>Abducens nerve palsy; painless proptosis?</td>
<td>Solid</td>
<td>Lateral orbitotomy</td>
<td>Partial</td>
<td>Partial recovery</td>
<td>?</td>
<td></td>
</tr>
<tr>
<td>Rui Miguel et al.[1]</td>
<td>42/male</td>
<td>Abducens nerve palsy; painful proptosis?</td>
<td>2.2</td>
<td>Cystic</td>
<td>Lateral orbitotomy</td>
<td>Total</td>
<td>Total recovery</td>
<td>?</td>
</tr>
<tr>
<td>Feichtinger et al.[8]</td>
<td>53/female</td>
<td>Abducens nerve paresis, optic atrophy</td>
<td>4.5x2x1.5</td>
<td>Solid</td>
<td>Lateral orbitotomy</td>
<td>Simple</td>
<td>Partial recovery</td>
<td>?</td>
</tr>
<tr>
<td>Present case</td>
<td>32/female</td>
<td>Pain in right eye</td>
<td>2.1</td>
<td>Solid</td>
<td>Superior orbitotomy</td>
<td>Total</td>
<td>Preserved</td>
<td>8 months</td>
</tr>
</tbody>
</table>
Figure 1: Magnetic resonance imaging T2 iso intense lobulated intraconal lesion displacing right optic nerve medially. The lesion is seen extending from the superior orbital fissure and closely abutting the right lateral rectus muscle

Figure 2: Magnetic resonance imaging coronal sequence showing T1 hypo and T2 iso intense lobulated intraconal lesion occupying the right orbital Apex with superomedial displacement of the optic nerve

Figure 3: Magnetic resonance imaging postcontrast axial (a) and sagittal (b) sequences showing intense enhancement

Figure 4: (a) Microphotograph showing a circumscribed thinly capsulated spindle cell neoplasm (H and E, ×40). (b) Microphotograph showing cellular Antoni areas with Verocay bodies (H and E, ×100). (c) Microphotograph showing hypocellular Antoni B area with a loose, vacuolated stroma (H and E, ×100). (d) Microphotograph showing nuclear palisading constituting the Verocay bodies (H and E, ×200)

Figure 5: Postoperative photograph of the patient showing synchronous complete range of extraocular movements of both eyes

Figure 6: Postoperative computed tomography scan showing complete excision of the tumor, there was no radiological evidence of bone formation in the polycaprolactone scaffold which is known to have osteo-inductive properties

and function as we could visualize the tumor attachment at myoneural junction, and safe and complete tumor excision was possible. With the availability of PCL scaffolds the orbital roof can be readily reformed.

Gamma knife surgery (GKS) has been tried in four patients of intraorbital schwannoma. After GKS, the patient treated for an orbitocavernous schwannoma experienced a significant deterioration in vision, temporary blindness in one eye, and late development of permanent abducens nerve palsy, which were
seemingly caused by compression of neurovascular structures within the annulus of Zinn during a temporary increase in the lesion's volume after irradiation. The cisternocavernous schwannoma underwent cystic degeneration 2 years after GKS, and the patient developed diplopia. Purely intracavernous neoplasms in general followed uneventful posttreatment courses. Imaging studies showed temporary enlargement of all tumors during the 1st posttreatment year, but thereafter, there was a trend toward reduction in volume and none of the neoplasms displayed regrowth. GKS controls the growth of abducens nerve schwannomas and may be effectively used to manage intracavernous neoplasms, but however, caution should be used in cases of dumbbell-shaped tumors, particularly those extending through the superior orbital fissure.

**Conclusion**

Though functional preservation of the abducens nerve is the rule, complete excision of the tumor should be the goal as residual tumor can potentially regrow, and GKS has discouraging outcome. Superior orbitotomy is an ideal approach as it gives direct access to nerve and muscle, which can be safely preserved while complete tumor excision is being achieved.

**Acknowledgment**

We thank our patient who agreed and gave consent for photographing and submitting the work for publication.

**References**