SUMMARY:

Forty-eight patients with intraorbital pathologies treated surgically and the literature are reviewed. Fifty per cent of the patients (28 male and 20 female) were between one and 20 years old. 26 cases had right and 22 left exophthalmus. 23 cases had 2nd, 3rd, 5th, and 6th nerve palsies. In 15 cases chemosis and in 13 cases pain (around the orbit) were observed. Pathological findings on plain x-ray were found in only 26 cases. In 33 (68.75%) cases tumoural and in 15 (31.25%) cases nontumoural pathology was found. Orbital unroofing, lateral orbitotomy, percutaneous needle puncture and embolisation were the treatments used.

KEY WORDS:
Orbit, orbital diseases, orbital tumours.

INTRODUCTION:

The orbita is a bony compartment which contains the globe, extraocular muscles, nerves, vessels, the lacrimal glands and adipose tissue. Conically shaped, with the apex directed medially, it has a volume of approximately 30cc. 6.5cc of which is the globe (1). Because of the organ it contains and its position, the orbita has an important function and is an aesthetically important part of the body. The orbita and its contents, the surrounding tissues or a metastatic process can be the cause of pathology. In addition infections and trauma can cause orbital pathologies. Because of its structure and the various sources of pathologies involved, the authors discuss the best treatment and operative approach.

Since there are many kinds of treatment, such as systemic and/or focal medical treatment, chemotherapy, or radiotherapy orbital pathology usually needs multidisciplinary evaluation.

Orbital deformation or loss of vision are the most common symptoms caused by a mass lesion and the diagnosis must be made after detailed physical and radiological examination. If surgery is indicated, a surgical plan must be made carefully, considering possible secondary defects.

Our experience of orbital pathologies, evaluated and surgically treated in the University of Istanbul Cerrahpaşa Medical Faculty, has been reviewed and compared with the literature.

MATERIALS AND METHODS:

Forty-eight patients with identified intraorbital pathology hospitalized between 1982 and 1989 were studied retrospectively. All the cases were treated surgically in the University of Istanbul Cerrahpaşa Medical Faculty Department of Neurosurgery. Clinical features (age, neurological and neuroophthalmological findings, site of the pathology, etc.), radiological findings, histopathology, surgical technique and the survival were discussed and results were compared with the literature.

RESULTS:

Twenty-eight cases were male (58.3%) and 20 were female (41.7%) (Fig: 1). Fifty per cent of the patients were between one and 20 years of age (Fig: 2). The childhood group included 17 (25.4%) cases. The youngest patient was two and the oldest 72 years of age. The mean age was 30.6. Varying degrees of proptosis were observed in all patients, 26 had proptosis in the right eye and 22 in the left. The direction of the proptosis depended on
In 27 patients (56%) plain skull plus orbita x-rays revealed pathological findings (Tab: 2). Angiography was performed in 31 patients but pathological signs were found only in 5 (10%) (3 blushing, 2 vascular anomalies). CT was performed in all cases and pathology was seen in all.

**TABLE : 2 PLAIN X-ray FINDINGS**

<table>
<thead>
<tr>
<th>FINDING</th>
<th>No of CASES</th>
<th>PATHOLOGY</th>
</tr>
</thead>
<tbody>
<tr>
<td>High density</td>
<td>8</td>
<td>Meningioma, Osteoma, Muco-sel, Haemangioma, Fibrous Dysplasia.</td>
</tr>
<tr>
<td>Lytic Lesion</td>
<td>4</td>
<td>Epidermoid, Meningioma, Infection.</td>
</tr>
<tr>
<td>Hyperostosis</td>
<td>4</td>
<td>Osteoma. Fibrous Dysplasia.</td>
</tr>
<tr>
<td>Fracture</td>
<td>2</td>
<td>Blow out Fracture, Carotico-cavernous Fistula.</td>
</tr>
<tr>
<td>Destruction</td>
<td>8</td>
<td>Meningioma, Hydatid Cyst, Adenoma, Abscess, Encephalocele, Fibrous Dysplasia, Rhabdomyosarcoma, Adenocarcinoma Metastasis.</td>
</tr>
</tbody>
</table>

Pathology was intracanal in 20 (40%) cases and extracanal in 28 (60%). Pathology was observed superiorly in 20 cases laterally in 8, medially in 4, inferiorly in 2 and apex in 14 according to the surgical anatomy.

In 33 cases (68.75%) tumoral and in 15 (31.25%) nontumoural pathology was found. 16 neural and 8 osseous pathologies formed the tumoural group. The 15 nontumoural pathologies included 5 infectious, 4 trauma and 6 different pathologies (Fig.5). In the childhood group there were 12 tumoral and five nontumoural cases (Table: 3.4).

The transcranial approach was the most used for the treatment of orbital pathologies (44 cases). Lateral orbitotomy in one case and embolisation techniques in 2 cases were preferred in our series.
PATHOLOGIES that originate in or invade the orbit produce symptoms and signs by compression, infiltration and/or infarction of the orbital structures (7,10,15). The most common clinical findings caused by orbital pathologies are; proptosis, optic neuropathy, ocular motor nerve paresis, pain, pupillary abnormalities and chemosis. In our series the most common features were proptosis, cranial nerve palsy, pain and chemosis. We found no isolated pupillary abnormalities, probably because tumours that produce pupillary sympathetic or parasympathetic nerve palsy are usually so extensive that they produce oculomotor nerve palsy with pupillary involvement as well. Thus, if there is damage to either the sympathetic or parasympathetic fibres it is usually masked by the oculomotor nerve palsy.

Correct diagnosis of orbital pathology has increased since the use of high resolution computed tomography (6,10,19,20). CT revealed pathology in all the cases in our series.

Pathological findings on direct x-rays were found in only 26 cases (56%). These findings were definite especially in the osseous tumours and meningiomas and our percentage was higher than that in Llyod’s series of 1070 cases, when a definite diagnosis was made in 21 percent (6,10,11).

Angiography which needs techniques such as selective catheterization, magnification, or subtraction is not often used in the evaluation of primary orbital pathology. Its principal role is in detecting small vascular tumours such as meningioma and haemangioma (6,8,10). Showing vascular displacement is very difficult since it is generally small, and is why only 10 percent of pathological findings were identiﬁed.

**DISCUSSION**

The characteristics of series of intraorbital pathologies vary due to racial characteristics, socioeconomic levels and the specialties of the authors. Definitions of pathological types differ in different series, but there are common points in typing the pathological groups (tumoural or nontumoural). Generally tumours are the most common type of orbital pathology, neurogenic tumours, haemangiomas, metastatic tumours, tumours of the lacrimal gland and osseous tumors being the most common (8,10,12,14,16,22). On the other hand some authors (4,13,18) report inflammatory pathologies as the most common, and in this group pseudotumours and endocrine exophthalmia have a high percentage. We confirmed in our series that 68.75% were tumoural and 31.25% nontumoural. 48% of the tumoural pathologies were of neural origin, and the others were mostly osseous. In some series lymphomas were the most common tumoural groups after haemangioma and optic glioma. In our series how ever we found no lymphomas (10), and about 17 per cent were nontumoural.

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**FIG. 5: ORBITAL PATHOLOGIES.**

**TABLE 3: TUMORAL ORBITAL PATHOLOGIES**

<table>
<thead>
<tr>
<th>Neural</th>
<th>Osseous</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optik glioma</td>
<td>Fibrous dysplasia</td>
<td>Metastasis</td>
</tr>
<tr>
<td>Meningioma</td>
<td>Osteoma</td>
<td>Adenoma</td>
</tr>
<tr>
<td>Neurofibroma</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 4: NON-TUMORAL ORBITAL PATHOLOGIES**

<table>
<thead>
<tr>
<th>Infection</th>
<th>Trauma</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydatid Cyst</td>
<td>Blow out fracture</td>
<td>Mucocel</td>
</tr>
<tr>
<td>Abscess</td>
<td>Caroticocavernous Pseudotumor</td>
<td></td>
</tr>
<tr>
<td>Infection</td>
<td>Fistula</td>
<td>Encephalocele</td>
</tr>
</tbody>
</table>

pathologies were removed totally in 28 cases, subtotally in 12. and only optic canal decompression was performed in two cases. In one patient with widespread rhabdomyosarcom orbital exenteration was performed at the same operation.

After surgical treatment 38 cases (79%) completely recovered and six remained the same. One optic sheet meningioma, one haemangioma and two osteomas localized around the optic canal became worse.

Decrease of visual acuity, chemosis, infection, paralysis of the extraocular musder were seen in 10 patients in the early postoperative period. These findings disappeared within a few days with drug therapy.
found in our 31 patients in which angiography was performed (2 meningioma, 1 hemangioma, 2 carotico-cavernous fistulas). The direction of orbital veins is more stable than arterial directions, so orbital venography is more significant for showing intraorbital pathology than arteriography (6,8,10).

The orbita can be divided into 4 parts (1): 1) Superiostial space, 2) Extraconal space, 3) Intraconal space, 4) Tenon space. The orbita has 5 sides. all the orbital walls may be used to reach the pathology. Generally there are four different types of approach (1,2): 1) Superior surgical approach, through the superior wall used during transcranial or transfrontal orbitotomy, 2) Lateral surgical approaches, external lateral or temporal orbitotomy is access through the temporal fossa (Krönlein technique or its variations) (Fig.6), 3) The anterior surgical approach can be performed through the eyelid or the conjunctiva (transethmoidal orbitotomy, transmaxillary orbitotomy) (Fig.7), 4) Percutaneous needle puncture. The route of access to the orbit is chosen according to the pathological condition (for example size, vascularisation, probable character of pathology) and surgeon’s specialisation (experience, working team and knowledge about of the orbital pathology) (1.2,3,7,8,12,13,21). It is possible to make a true diagnosis during the preoperative period with delicate neurological examination and using new diagnostic techniques for optic nerve or sheet pathologies (1.8,10,18,19,20). This evaluation is important for choosing the surgical approach. The transcranial approach was used in 44 cases, lateral orbitotomy in one, percutaneous needle puncture in one case and embolisation in two cases (carotico-cavernous fistula).

Optic gliomas become manifest during the first decades of life and incidence increases with neurofibromatosis. They are usually low grade tumours (grade 1–2), with 0.6-1.2 percent incidence among cases of unilateral proptosis (3,8,10,15,17). Optic glioma may be in the prechiasmatic region in 23-36 percent in the chiasma, tractus region (17). Adult optic gliomas are high grade astrocytomas, and progress more rapidly than childhood astrocytomas (6,8).

Fig.6: Variation of lateral orbitotomy.

Therapeutic methods depend on the chiasmal infiltration. The authors Llyod, McKeran, Hoyd, Donol, Robert who have large series, recommend extirpation of the glioma by cutting the optic nerve in front of the chiasma and just behind the globe if the tumours is in the prechiasmatic region and does not infiltrate the chiasma in patients whose vision is impaired and who have progressive proptosis (3,4,9,11,12,13,17). They recommend only biops and RT if chiasmal infiltration is present. Our general practice is to evaluate the vision and CT estimation of the tumor size if the glioma is intraorbital and confined to one nerve. If vision is seriously impaired and proptosis is progressive we use radical surgical techniques to eliminate any change of chiasmal spread. Four patients on whom we performed total extirpation, had 0-1m vision before the operation and to date none has had tumoural recurrence.

The efficiency of RT is suspect in the literature for optic nerve or sheet tumours, but some authors say it is 50 percent useful for patients with tumours sited posteriorly (chiasma, tractus) and those in whom partial tumoural resection has been performed (17).

There is some controversy about repair of the orbital roof after unroofing (1.2,5,13). When exposure of the orbital cavity has been extensive, or when a subsequent operation such as orbital exenteration is planned, repair of the orbital roof is recommended. The repair can be done either with a sheet of plastic polymer or with a bone graft. We could not repair the orbital roof after orbital unroofing in our cases but did not observe any problems during the early and late stages of the operation.

We observed some complications, such as chemosis, ptosis, extraocular motor nerve palsies, impairment
of vision, in 10 patients during the early postoperative period which improved within days. The signs regressed with corticosteroid therapy and were probably temporary paralysis of the levator palpabra and superior rectus muscles due to surgical manipulation during the transcranial approach reported in the literature (2,4).

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REFERENCES: