TECTAL GLIOBLASTOMA

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ABSTRACT - Brain stem gliomas are a heterogeneous group of neoplasms arising mostly in paediatric patients. Tectal plate gliomas represent a particular type of brain stem tumours usually with a benign, indolent clinical course, presenting with signs of raised intracranial hypertension due to supra-tentorial hydrocephalus caused by aqueductal stenosis. Seldom high-grade lesions arise in this location with tremendous therapeutic implications. When a malignant tumour is clinically and radiographically suspected a biopsy should be performed to obtain histological confirmation. Treatment is then planned in a case-by-case basis. We present the case of a glioblastoma of the tectal plate in a 22 years-old woman operated upon by a supracerebellar-infratentorial approach.

KEY WORDS: brain stem, tectal plate glioma, glioblastoma.

Brainstem tumours are usually described as paediatric neoplasms. In adults they are not so well known and account for less than 2% of gliomas¹. The peak of incidence is the first decade in children and third and fourth decades in adults². There are no significant clinical differences between the two groups with mean duration of symptoms being usually longer in adults. Survival may be significantly shorter in children with brain stem gliomas when compared with older patients. Histopathological analysis revealed grade II astrocytomas as the majority of gliomas in children and adults. The grade of brainstem glioma is a significant predictor factor for survival in adults, but in children it may not correlate with outcome⁶. Low-grade gliomas tend to occur in the upper brain stem and high-grade tumours in its caudal portion. Tectal plate gliomas are a distinct group of midbrain tumours, usually with a benign course¹. Periaqueductal location and absence of contrast enhancement and long periods of stability are classic features³. They have a better prognosis because of their non-infiltrative pattern and slow growth rate.

We report an illustrative case.

CASE

A 22 years-old woman, presented with insidious and low intensity generalized headache, together with paresthesia of the right body for the last month. MRI revealed an extensive, contrast-enhancing lesion located in the tectal plate, with supratentorial hydrocephalus. A neuroendoscopic ventriculostomy and biopsy were performed at another hospital. Histopathological analysis was consistent with glioblastoma (grade IV WHO astrocytoma). She did not undergo any complementary therapies.

At the time of consultation in our institution, six months...
later, she presented with multidirectional nystagmus, visual impairment (greater on the right) and epicritic (thermo-al-gic) incomplete hemi-sensory loss on the right side, without any other neurological deficits. Pre-operative MRI showed an exophytic lesion in the tectal plate (Fig 1).

A ventricular-peritoneal shunt was placed and four weeks later a partial resection of the tumour was achieved by a supracerebellar-infratentorial mid-line approach. A complete third pair palsy on the right was the only side effect of the surgery (Figs 2 and 3).

Post-operative MRI showed partial removal of the tumour without complications (Fig 4).

Histopathology continued to reveal a grade IV WHO astrocytoma.

She underwent complementary treatment with both radiotherapy and chemotherapy (temozolamide) and, six months after the procedure, she is clinically and neurologically stable.

**DISCUSSION**

Brainstem glioma, pilocytic astrocytoma and medulloblastoma are the most frequent infratentorial tumours in patients under 18 years representing 10-30% of brain tumours in children. They are usually infiltrative lesions and only a small number (dorsal exophytic) have a favourable prognosis. They occur mostly in childhood and adolescence (77% in ages below 20 years), representing 1% dos tumours in adults.

Midbrain tumours are a heterogeneous group of neoplasms with variable clinical and radiological features, relating with the location and tumour histology. They occur in the tectal plate, tegmentum, invading the pons or cerebral aqueduct. Sometimes
they represent midbrain invasion by tumours of adjacent regions, namely pineal and thalami.

Tectal gliomas are in majority low-grade astrocytomas, considered a “benign” sub-group of brain stem gliomas. They represent approximately 10% of brain stem gliomas in children and 6% of paediatric brain tumours surgically treated.

Computhorized tomography (CT) reveals the hydrocephalous but may not be able to detect tectal plate tumours in up to 50% of patients. Calcifications are seen in 9-25% of cases. Magnetic resonance imaging (MRI) is the chosen exam for diagnosis and follow up of tumours in this location. It allows a precise evaluation of the growth pattern and correct preoperative diagnosis in most of cases. Gadolinium enhancement, calcifications, cysts e exophytic nature are observed in both low and high-grade gliomas. They are typically isointense in T1WI and iso or hiperintense in T2WI. Enhancement after endovenous contrast enclosures an undefined pathological significance. In the case of intrinsic tectal tumours, low-grade astrocytoma is the probable diagnosis. Differential diagnosis of exophytic tectal tumours includes pineal neoplasms, requiring histological verification.

Since biopsy is not performed upon many lesions, a precise statistic analysis is not possible.

They are mainly astrocytomas (pilocytic astrocytoma, WHO II diffuse astrocytoma, anaplastic astrocytoma, high-grade astrocytoma) but other lesions have been identified (oligodendrogloma e oligoastrocytoma, WHO II ependymoma, ganglioglioma, medulloblastoma, primitive neuroectodermal tumours, dis-embryoblastic neuroepithelial tumours, metastasis, melanoma, lipoma, cavernoma, abscess and periaque ductal gliosis).

Many neurosurgeons perform a stereotactic biopsy to obtain histopathological confirmation of a low-grade tumour and only then the treatment is planned.

Clinical presentation with signs of raised intracranial pressure due to cerebral aqueduct compression resulting in supratentorial hydrocephalous is the most common clinical feature affecting all patients in some series. Focal neurological findings are less frequent (as diplopia, visual field defects, nystagmus, Parinaud syndrome, seizures) and usually revert after correction of the hydrocephalous.

It is not universally accepted that lesions with radiographic progression need to be treated. Paediatric tectal plate gliomas are usually low-grade tumours that can be managed conservatively even in the presence of radiographic enlargement, reserving radiotherapy and chemotherapy for clinical progression which is described in 15-25% of cases. It is even more advantageous to observe these patients in order to avoid radiation therapy and chemotherapy induced neurodevelopmental and endocrinial injury to the developing brain.

The initial treatment is directed to correction of hydrocephalous. Ventricular-peritoneal shunt placement has good long-term results, if no dysfunction is verified. Third endoscopic ventriculostomy suppresses the need for shunt placement and a biopsy can be performed through an enlarged foramen of Monro. It allows resolution of signs and symptoms and the return of the ventricular system to its normal size. It is the procedure of choice for paediatric patients. Endoscopic aqueductoplasty with flexible systems (stent based or not) may be an option for some cases, but its long-term results are unknown.
Due to its indolent course, open surgery is not usually indicated for low grade tumours. However, if a malignant, secondary or vascular lesion is clinically and radiographically suspected a microsurgical procedure should be performed.

Simple stereotactic aspiration of cystic brain stem gliomas is not an effective treatment strategy, because they will frequently recur leading to progressive neurological deficit. When combined with stereotactical placement of intra-cyst catheters, intracavitary irradiation with radioactive solutions, external radiotherapy and chemotherapy, it may allow cyst control without permanent morbidity or mortality.

Resection or open biopsy of tumours in this location can be achieved by a supracerebellar-infratentorial or suboccipital-transtentorial approach, with the extent of removal being wider at the level of the superior colliculi and limited at the inferior colliculi due to high auditory risk. Parinaud syndrome is one of the most frequent surgical complications. Auditory hallucinations and acoustic neglect syndrome can also occur. Despite of that, tectal plate region is a safer surgical field than the ventral midbrain.

Early and middle-latency brain stem auditory evoked potentials should be used for functional brainstem evaluation and definition of resection margins during tectal plate surgical procedures.

Stereotactic radiosurgery can be employed on tumour progression but due to radiation side effects the dosage is limited.

Optimal treatment of tectal plate gliomas is still to be determined. The role of different treatment modalities is unclear and universally accepted guidelines are still to be proposed. Serial neurological / clinical observations and MRI scans each 6-12 months is an option.

Patients with well-differentiated brainstem gliomas may be cured by microsurgical resection.

Like in all high grade gliomas, resection of the tumour instead of biopsy, age equal or less than 60 years and a Karnofsky scale of 70 or greater are all correlated with better outcome.

Neuroanatomy based craniotomy for tumour resection is the mainstream of treatment currently available if it can be done safely, without further neurological deficits. However, surgical resection alone does not cure malignant brain tumours unless it is coupled with other treatment modalities addressed to the diffuse nature of these lesions, like chemotherapy and/or immunotherapy.

In high-grade gliomas, like the case presented, partial resection may prolong survival and facilitate subsequent complementary therapeutics.

REFERENCES