Gliomatosis cerebri in a 10-year-old male patient

Sir,

Gliomatosis cerebri is a rare, diffuse form of glioma. The World Health Organization describes it as a diffuse glial tumor infiltrating at least two cerebral lobes, often extending to infratentorial structures and preserving the anatomic architecture. The largest series to date describing pediatric cases is the one by Armstrong et al.,[1] that included 13 patients in a 20 years-experience, though most reports include no more than 3 patients.[2,3] We would like to describe the case of a 10-year-old male patient who presented to our institution with a 5-month history of symptoms and a relatively aggressive course.

A 10-year-old male patient was referred to our institution after a 5-month course of neuropsychiatric and neurologic symptoms consisting of behavioral changes (anhedonia, depression) and loss of strength in the right limbs and face. An magnetic resonance imaging (MRI) showed a diffuse T2-hyperintense bilateral lesion involving the cerebral hemispheres. Initially, he was treated in another hospital with pulse therapy with methylprednisolone for an acute demyelinating disease (the patient had a history of concomitant amygdalitis), but with no response.

The initial neurologic examination revealed a right supranuclear facial palsy, Grade 2 proximal and Grade 3 distal strength in the upper limbs and Grade 4 strength in the upper and lower left limbs; vivid and symmetric reflexes, Babinski sign on the right and left dysmetria. Blood and cerebrospinal fluid studies were normal. MRI showed a diffuse lesion affecting the thalamus, basal nuclei and white substance bilaterally, hyperintense in T1-weighted and T2-weighted imaging and with no contrast enhancement. Spectroscopy showed a choline peak [Figure 1].

The patient was submitted to a left parietal bur hole stereotactic biopsy, with no complications. Perioperative anatomopathological examination showed that the specimens were indeed correctly obtained. Definitive anatomopathological analysis revealed a Grade 3 astrocytoma [Figure 2a and b] and immunohistochemistry showed positivity for glial fibrillary acidic protein and the proliferation index of 20% [Figure 2c and d].

The patient was started on chemotherapy with temozolomide and concomitant radiotherapy (60 Gray). A new MRI, 30 days later, showed further growth of the lesion and now contrast enhancement was noted [Figure 3]. There was a progressive neurological deterioration and parents decided to stop treatment before the end of radiotherapy.

Histological analysis of samples usually reveals a Grade 2 or 3 astrocytoma. Treatment remains unclear. There is a marked better survival for patients submitted to treatment.[1] Radiotherapy is one of the most popular treatment strategies used in the literature, although concomitant chemotherapy is being increasingly used, and results show a potential increase

Figure 1: T2-weighted image (a) and T1-weighted image (b) revealing a diffuse hyperintense lesion involving the white matter and the diencephalon. Below, a spectroscopy study showing a choline peak (c)
Considering only the pediatric population, the median survival is 27 months.\(^1\) Taking into consideration all the above, we stress the need for early diagnosis (mostly depending upon brain biopsy) in order to establish the most adequate treatment – since gliomatosis cerebri still carries a poor prognosis.

**Financial support and sponsorship**
Nil.

**Conflicts of interest**
There are no conflicts of interest.

**References**


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**How to cite this article:** Strapasson A, Antunes AC, Oliveira F, Oppitz PP. Gliomatosis cerebri in a 10-year-old male patient. Asian J Neurosurg 2017;12:336-7.