Complete surgical resection of high-grade astroblastoma with long time survival: case report and review of the literature

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Summary

Astroblastoma is a rare glial neoplasm of unknown origin and uncertain prognosis. It usually presents in young adults as a well circumscribed hemispheric mass, often associated with a cystic component. The histological features of astroblastoma are the presence of typical astroblastic perivascular pseudorosettes and perivascular hyalinization. Two different subtypes of astroblastoma have been defined based upon histological characteristics. Prognosis, however, sometimes is in contradiction with the pathological appearance and seems to be more closely related to the grade of surgical resection. We present a new case of a patient with a high-grade astroblastoma with a long survival time, in whom complete surgical resection was confirmed by an early postoperative MRI.


Astroblastoma de alto grado con supervivencia prolongada tras resección completa: descripción de un caso y revisión de la literatura

Resumen

El astroblastoma es un tumor glial infrecuente de origen desconocido y pronóstico incierto. Habitualmente se presenta en adultos como una masa hemisférica bien definida, presentando a menudo un componente quístico. Las características histopatológicas de este tumor incluyen la presencia de pseudorrosas astrocíticas y la hialinización perivascular. Se han distinguido dos tipos de astroblastoma en relación a la malignidad histológica. Sin embargo, el pronóstico en ocasiones no guarda relación directa con el tipo histológico y parece más dependiente del grado de resección quirúrgica. Presentamos el caso de una paciente con un astroblastoma de alto grado con supervivencia prolongada, en el cual la resección quirúrgica completa se demostró mediante resonancia magnética postoperatoria precoz.


Introduction

Astroblastomas are glial neoplasms of unknown origin characterized by a typical perivascular pattern of GFAP-positive astrocytic cells with broad non-tapering processes radiating towards a central blood vessel. Since a similar histological pattern may occur in low- and high-grade astrocytomas as well as in glioblastomas, the term astroblastoma should be restricted to those rare tumors in which it prevails throughout a well-demarcated lesion that does not contain foci of conventional or infiltrative astrocytoma or ependymoma. Prognosis is still uncertain. Complete surgical resection has been sometimes associated with long survival times even in cases showing histological features of malignancy.

Case report

A 42-year-old woman presented with a 3-month history of headache and a recent episode of a complex partial seizure with secondary generalization. Physical and neurological examination were both normal.

Cranial CT showed a left frontal cortico-subcortical lesion, isodense to the brain parenchyma, with heterogeneous enhancement after intravenous contrast administration. Cranial MRI also revealed heterogeneous areas inside the lesion which showed well-defined borders. There was

mild edema of the surrounding brain and little mass effect on the ipsilateral ventricle.

The preoperative diagnosis was high grade glioma. At surgery, a well defined cortico-subcortical lesion with small cysts in its deepest part was found. Apparently, it was completely resected and an early cranial MRI, performed within 24 hours of operation, confirmed the absence of tumoral rests.

Histopathological examination showed an extensive perivascular pseudorosette pattern of astrocytic cells with broad processes radiating towards central vessels; small foci of hyalinization were frequent. In some areas cellularity was increased with pleomorphism, high mitotic rate, small necrotic areas with perinecrotic palisading and hypertrophy and hyperplasia of vascular endothelia. Tumoral cells were GFAP, vimentin and s-100 positive, preferentially in the perivascular processes.

The postoperative course was uneventful. The patient underwent adjuvant radiation therapy. One month after surgery the patient had returned to her previous job as a teacher and eighteen months later she remains asymptomatic and both MRI and cranial SPECT show no recurrence.

Discussion

Astroblastoma is a rare type of glial tumor that was firstly described by Bailey and Buey in 1930. Although its existence has been questioned by some investigators, it is now recognized as a distinct clinicopathological entity, probably representing less than 0.5% of gliomas. It usually appears in young-adult patients, although congenital cases have also been described. No sex preference has been observed.

Clinical symptoms at presentation depend on locali
zation, size and mass effect of the neoplasm. Headache and seizures are often present. The cerebral hemispheres are most commonly affected, but astroblastoma may also develop in other places of the central nervous system, such as the corpus callosum, brainstem or cauda equina.

Radiologically, astroblastoma usually appears as an hemispheric, well circumscribed, often cystic mass, located peripherally near or at the surface of the brain. Thus, differential diagnosis with extraaxial neoplasms is sometimes necessary. Astroblastoma tends to be isodense to brain parenchyma on the plain CT scan.

Califications may be seen. The solid component is contrast enhancing. On MRI, astroblastoma appears hypointense in T1-weighted sequences and hyperintense on both proton-density and T2-weighted sequences. Brain edema, if present, is mild. MR spectroscopy findings documented are consistent with those observed in other brain neoplasms and are particularly similar to those of astrocytoma.

According to the WHO, astroblastoma belongs to the category of neuroepithelial tumors of unknown origin. Rubinstein and Herman observed a close ultrastructural and immunohistochemical similarity between astroblastoma cells in culture and the tanyocyte, which is a cell integral to the ependyma of submammalian species with transitional features between the astrocyte and the ependymal cell. It has been suggested that a similar cell may also occur transiently during normal human embryogenesis and that astroblastoma may derive from abnormally persisting examples of such embryonal precursor cells. Therefore, despite its designation, astroblastomas are not considered to be embryonal neoplasms, but rather of astrocytic lineage if their characteristics are taken into account.

Concerning the cytological architecture of astroblastoma, Mierau et al demonstrated two morphologically distinct cell types displaying an unusual organizational relationship: more primitive cells appeared nesting within the cytoplasm of the differentiated cells. The authors hypothesized that the more differentiated cells served as "nurse" cells for the maturing population.

Histologically, astroblastoma is characterized by a typical perivascular pseudorosette pattern of GFAP-positive cells with broad, non-tapering processes radiating towards a central blood vessel. A component of perivascular hyaline is usually present. Focal astroblastic features are nonspecific and can be seen in low and high grade astrocytic tumours. As a consequence, the term astroblastoma should be reserved for those rare tumours in which the pattern prevails throughout a well demarcated lesion. Interestingly, a case of astroblastoma of the pure type has also been reported.

In a pathological study of 23 astroblastomas with a postoperative follow-up in 13 patients, Bonnin and Rubinstein observed two distinct histological types: 1) a lowgrade type, in which a better differentiated pattern was apparent, and a favourable postoperative prognosis may be expected in a fairly large proportion of patients; and 2) a high-grade type, showing more anaplastic features and in which the length of postoperative survival was usually shorter. Focal or multifocal regions of high cellularity, anaplastic nuclear features, elevated mitotic indices (>5 per 10 HPF), vascular proliferation and necrosis with pseudopalisading correspond to this high-grade group.

Immunohistochemically astroblastomas are immunoreactive for GFAP, S-100 protein and vimentin. The majority also display at least a focal cytoplasmic immunoreactivity for EMA. We have previously reported a case of low-grade astroblastoma in which due to the scarce positivity with GFAP, in contrast with the abundance of intermediate filaments in electron microscopy, along with immunohistochemical and ultrastructural findings, lead us to conclude that the filaments seen in tumor cells were mainly vimentin filaments.

Recently, molecular analysis has been applied to the study of chromosomal abnormalities in astroblastoma. Brat et al, using comparative genomic hybridization, detected chromosomal abnormalities in every case studied, the gain of 20q being the most frequent. Although the spectrum of chromosomal alterations indicated that astroblastoma may have its own genetic profile, none of the detected anomalies were specific for malignancy. Shuangshoti et al using microsatellite markers, showed loss of heterozygosity (LOH) at the D19S412 locus on the long arm of chromosome. This observation was compatible with the loss of a tumour suppressor gene in this region. This oncogene is supposed to play a role in the tumoral genesis.

The prognosis of astroblastoma remains to be determined. In the absence of sufficient clinicopathological data, it has been decided not to establish a WHO grade at the present moment. Low grade astroblastomas seem to have a better prognosis than the high grade ones. Occasionally, however, in the presence of histological features strongly indicative of malignancy, the clinical follow-up may be in contradiction with the microscopical appearances. This is in contrast to astrocytomas, in which there is extensive concordant evidence that clinical behavior is closely correlated with histopathological features. Long term survival likely depends on numerous factors including tumor location, extent of resection and response to adjuvant therapy. In our case, complete surgical resection has probably played a major role in the long tumor-free survival of the patient. However, and given the histological malignancy of the neoplasm, postoperative adjuvant therapy was associated in this patient. Although astroblastoma appears to be radiosensitive, no definite results concerning chemotherapy are available.
The optimal management of astroblastoma remains to be established. As mentioned above, radical removal is important and given its peripheral location and well circumscribed aspect, it seems a goal easier to achieve than in other gliomas. In low-grade astroblastomas radical surgery may be the only necessary treatment, although cases of early recurrence have been documented. In such instances, re-operation and adjuvant radiotherapy are the better options though in cases of high-grade astroblastomas a complete resection may be also curative. The lack of enough information about the natural history favours the use of adjuvant radiotherapy from the beginning.

Early MRI postoperative MRI is helpful in demonstrating the absence of residual tumor after surgery that could be otherwise misinterpreted as tumor recurrence in ulceror studies. Therefore it seems very useful in helping to determine the influence of radical resection in the natural history, prognosis and best management of patients with astroblastoma. However, further studies are still necessary to define the real behavior of astroblastoma and determine its optimal management.

References