Primary diffuse leptomeningeal gliomatosis predominantly affecting the spinal cord: case report and review of the literature

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Abstract

Primary leptomeningeal gliomatosis is a rare, fatal neoplastic syndrome. A 71 year old man is reported on, who after a 2 month history of back stiffness, epigastric pain, and weight loss developed visual blurring. Cranial CT and MRI studies showed no leptomeningeal enhancement. Examination of CSF 10 weeks premortem showed an increase in protein and decrease in glucose but no malignant cells. He became increasingly confused and repeated CSF examination showed inflammation and a few suspicious cells but no definitive evidence of neoplasia. He died 7 months after onset of his initial symptoms. At postmortem meningeval whitening was seen at the base of the brain and over the spinal cord. Histology disclosed diffuse leptomeningeal gliomatosis (GFAP positive, cytokeratin negative) over the brain, optic nerves, and spinal cord without parenchymal involvement. No tumour was found in internal organs. The diagnosis of primary leptomeningeal gliomatosis was not evident after cranial CT and MRI and CSF examination premortem. Suspected cases need MRI scanning of the entire neuraxis and meningeal biopsy.

Keywords: leptomeninges; nervous system; tumour; necropsy; glioma

Primary diffuse meningeal gliomatosis (PDLG) is a rare, fatal condition characterised by widespread infiltration of the meninges by tumour apparently arising from heterotopic glial nests without evidence of tumour within the parenchyma of the brain or spinal cord. Heterotopic glial nests occur in the subarachnoid space in about 1% of unselected necropsies1—most often affecting the medulla (57%)—with a higher incidence (25%) in patients with congenital malformations of the nervous system.2

At presentation, most patients complain of headache,3 with signs of raised intracranial pressure or meningism. Differential diagnoses include carcinomatous meningitis and autoim-
Electromyography and nerve conduction studies were normal. Further MR studies were planned, but he was very restless and it was considered unsafe for him to have a general anaesthetic. Lumbar puncture was performed repeatedly (some 16 times over the 6 weeks until his death). The CSF pressure remained high (50–80 cm), protein just slightly increased (maximum 0.85 mg/dl), glucose very low (<1 mmol/l), and the white cell count rose gradually from 0 to 16 cells/µl. Tuberculous and cryptococcal testing were negative. Treatment over this period was largely supportive. An empirical course of intravenous methylprednisolone had no clinical effect. He was on empirical quadruple antituberculous therapy when he died on day 4 of this treatment on 20 October 1998.

Necropsy

The scalp, skull, dura, and venous sinuses were normal. Apart from bronchopneumonia no significant abnormality was found outside the nervous system. The brain weighed 1740 g and the leptomeninges showed focal areas of whitening over the vertex and base of the spinal cord with thickening maximal (2.5×2×1 mm) at T4 (figure A and B).

Microscopical examination

Histology showed leptomeningeal infiltration by a diffuse infiltrating astrocytoma over the brain, spinal cord, and optic nerves. Spindle shaped tumour cells with fibrillary processes and pleomorphic, vesicular nuclei were seen extensively within the reticulin meshwork of the leptomeninges. Tumour cells (figure C) expressed GFAP (polyclonal antibody, DAKO, 1:200). The proliferation rate (proliferation marker Mib-1 or Ki-67, monoclonal antibody, Immunotech, 1:50) was focally up to 20% (tumour at T4) and varied in other areas down to 3%. Tumour cell nuclei were p53 negative (monoclonal antibody, Novo Castra, clone DO7, 1:50 and 1:20). There was no discrete focus of tumour within the parenchyma of the brain or spinal cord. This was therefore considered a case of primary diffuse leptomeningeal gliomatosis.

Discussion

As well as being a rare diagnosis the case illustrates the wide constellation of symptoms and signs that PDLG may manifest. This is the seventh reported case of PDLG predominantly affecting the cord. Of the 21 cases of PDLG reported since 1957, 18 were verified at necropsy.

An increased CSF protein combined with low CSF glucose, especially in the presence of a high CSF cell count and raised intracranial pressure, as in our case, indicates a possible neoplastic process, and has been noted in most of the reported cases of PDLG. The raised CSF pressure in our case was most likely due to tumour mediated obstruction of CSF drainage into spinal nerve roots, as the bulk of the tumour was spinal rather than cerebral and the arachnoid granulations were free of tumour. The increased CSF protein concentration may also have contributed to the high pressure gradient.

The signs in the present case did not initially suggest the need for spinal MRI examination. In retrospect that might have been helpful in obtaining a diagnosis although given the small size of the lesion at T4 at necropsy it seems likely that this would not have been enhanced even with MRI earlier in the illness; MRI enhancement of the meninges has been noted in only a few reports. In reported cases of PDLG without a clear premortem diagnosis (as in our patient) antituberculous treatment is almost always tried. Patients treated additionally with radiation or chemotherapy survive slightly longer than with drug treatment alone; one patient even improved and a further patient went into complete remission for 15 months. If the CSF showed atypical cells and the patient's therapy included irradiation remission or clinical improvement is reported (by contrast with treatment with chemotherapy alone). Survival is also associated with different factors such as the World Health Organisation (WHO) grade of the
tumour, complicating lesions such as infarcts (due to vascular compression by adventitial tumour mass), and the site of the lesion—for example, involvement of vital centres. The overall poor prognosis (comparable with gliomatosis cerebri) suggests a worse biological behaviour than predicted by WHO grade of the biopsy in many cases. 

As the bulk of the PDLG as in this case may be spinal rather than cerebral, spinal as well as cranial MRI should be considered in suspected cases. In PDLG as in gliomatosis cerebri, MRI is superior to CT in detecting diffusely infiltrating neoplastic astrocytes.

We thank Steve Toms and Michael Todd for photographic assistance.