Progressive leptomeningeal fibrosis: a clinico-pathological case report

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SUMMARY A female patient developed persistent facial pain beginning at age 19 years. Intermit-
tent motor and sensory disturbances referable to one hemisphere began nine years later and by
the age of 41 she had developed signs of increased intracranial pressure. Exploratory craniotomy
revealed replacement of the leptomeninges by thick, fibrous tissue. The histological appearance
was that of a chronic, benign and minimally infiltrative process with a mild, non-specific
inflammatory component, underlying cortical ischaemic changes, and white matter oedema. The
lesion resembled nodular fasciitis, a soft tissue process. No cause of the reactive fibrosis of the
meninges in this case is known.

Benign fibrous lesions occur commonly in the soft
tissues yet rarely in the CNS or meninges. The
reported cases of intracranial fibroblastic prolifera-
tive processes form a spectrum histopathologically,
and usually present as well defined discrete
tumours.1 Leptomeningeal fibrosis may follow cra-
nial irradiation2 and sub-arachnoid haemorrhage3
but does not appear to have been reported as an
idiopathic entity. We report a case of diffuse fibrosis
of the leptomeninges of the right hemisphere in an
otherwise healthy 41-year-old female who was
aware of progressive symptoms referable to that
hemisphere for more than 20 years.

Case report

The patient is a 41-year-old right handed female surveyor,
who first came to attention at the National Hospital, Queen Square in 1972. She had suffered from persistent
right facial pain in the first two divisions of the fifth cranial
nerve for over 20 years, at times associated with right-
sided headache. For four years she had experienced
episodes of numbness lasting several days, over the left
side of the body associated with vague clumsiness of the
left hand. There was one isolated episode of loss of con-
sciousness associated with a numb sensation in the left leg.
Neurologic examination and general physical examination
were negative. EEG and EMG were normal. A radio-
isolette brain scan showed an area of increased uptake in
the right parietal region, however four- vessel angiography
was normal. The patient was discharged with a presumpt-
ive diagnosis of migraine.

She presented next in 1976 with clumsiness of the left
arm starting acutely several weeks prior to admission and
slowly subsiding. In the interim period (1972–76), the
left-sided motor-sensory phenomena had subsided, but the
right facial pain had persisted. Neurologic examination
showed left-sided hyperreflexia and mild weakness. There
was no papilloedema. Lumbar puncture yielded clear col-
ourless fluid with an opening pressure of 140 mm water, no
cells and a normal biochemical profile. There was no
oligoclonal pattern. CT scan (fig 1A) showed a low density
area in the right parietal lobe which enhanced with con-
trast. No mass effect was appreciated. EEG demonstrated
a right anterior quadrantic disturbance; regional cerebral
blood flows were normal. Right carotid angiography
showed small abnormal tortuous vessels over the right
posterior frontal convexity, supplied mainly by a dilated
anterior middle meningeal artery. The angiographic pic-
ture was thought to be consistent with low grade glioma.
As the patient had minimal neurologic deficit and as the
lesion was ill-defined, surgical exploration was deferred,
and she was discharged to be followed as an out-patient.

The patient was re-admitted in 1981 for investigation.
The right-sided facial pain had remained constant although
the right-sided headache had increased in severity and
would at times wake her at night. About two weeks prior to
admission she had developed a slowly progressive left-
sided numbness and she continued to have intermittent
episodes of left-sided motor and sensory disturbances.
There were no other symptoms. Family and past medical
history were non-contributory. Over the previous 20 years
the patient had taken a number of drugs for relief of
headaches and facial pain, including aspirin, codeine phos-
phate, paracetamol, propoxyphene and benzodiazepines. Specifically, there was no history of methysergide ingestion.

Examination revealed an alert, intelligent woman with mild pyramidal weakness and hyper-reflexia on the left side. Plantar responses were flexor. There was mild blurring of the optic discs and fluorescein retinal angiography showed early papilloedema. Repeat CT scan (figs 1b, c) demonstrated extensive areas of low density in the right hemisphere, with marked cortical enhancement after contrast infusion. Bilateral carotid angiography disclosed diffuse swelling of the right cerebral hemisphere. The areas of abnormal circulation in the right posterior frontal region were persistent and the contribution of the meningeal circulation was more apparent than previously.

The patient was started on steroids with some clinical improvement and underwent right frontal craniotomy for biopsy and decompression. The surface of the brain was grossly abnormal and were covered in most areas with a 3–4 mm thick layer of firm, pinkish tissue which was lightly adherent to the overlying dura. Dilated meningeal arteries were seen in the dura adjacent to the thickened leptomeninges. In some areas where the brain was not covered by the abnormal tissue the arachnoid had a white discoloration and was thickened along some of the sulci.

Following operation the patient made an excellent recovery. Treatment with steroids continued for several weeks during which time there was a continuing improvement in her left sided weakness. She was then able to exist without steroid therapy for several months but approximately one year after her operation she was re-admitted with a short history of deterioration during which her hemiplegia had worsened. At the time of her arrival at hospital some spontaneous improvement had started. Treatment with steroids was started once again and continued for several weeks. Further CT scans performed during this recurrence of her symptoms at first showed increased oedema in the right hemisphere but later scans demonstrated resolution of this oedema. At the time of writing she is back at work with only a slight increase in her hemiplegia compared with that noted prior to her operation.

Pathology

The operative specimen received consisted of brain tissue 1.5–2.0 cm in thickness including the frontal pole and adjacent gyri with underlying white matter. Many blocks were prepared so that the whole specimen has been embedded in paraffin wax. A wide variety of staining methods were used. The description which follows applies to every block studied with only minor variations. There was generalised thickening of the leptomeninges varying from about 100 µm in thickness over some gyral convexities (fig 2) up to 1.5–2.0 mm in the sulci. In places the leptomeninges were cellular, composed of plump fibroblast-type cells (fig 3) lying in a loose reticulin-rich matrix and showing moderate numbers of mitotic figures. There were scattered clusters of lymphocytes, some around vessels, but no necrosis, granulomas or plasma cells. No organisms were identified with appropriate staining techniques for bacteria or fungi. In other places the meninges were densely collagenous with few cells, and here and there small vessels and collagen had an eusinophilic hyaline appearance (not positive for amyloid). Arterial and venous vessels were present within the thickened leptomeninges as well as endothelium-lined channels. Although vessels were numerous in some places, particularly in sulci, the tissue was not vascular enough to be considered angiomatosus and only occasional granules of haemosiderin pigment were present, too little to indicate previous bleeding. Vessels were surrounded by concentric layers of collagen and some seemed to be constricted by the dense surrounding tissue. Several vessels were seen which have a greatly narrowed lumen because of intimal thickening and smooth muscle proliferation suggesting previous occlusion and recanalisa-
Fig 2  Section showing meninges and underlying cerebral cortex at the convexity of a gyrus. There is marked thickening of the pia-arachnoid (arrow) which is here about 200 μm thick, densely collagenous with a scattering of lymphocytes. There are no granulomas. Immediately below the pia there is a broad (about 80 μm) band of glial fibres (*) and the molecular layer beneath shows reactive astrocytic gliosis. (Haematoxylin and eosin × 40.)

Fig 3  Section showing a cellular area of meninges. Fibroblasts and occasional lymphocytes lie in a loose matrix which, with appropriate staining, can be shown to be rich in reticulin fibres but contains little collagen. (Haematoxylin and eosin × 100.)

tion. There was no arteritis or phlebitis and no acute thrombosis. Perivascular fibrosis was seen around some vessels in the white matter (fig 4) and may represent an extension of the fibrous lesion along Virchow-Robin spaces.

The underlying brain showed changes which could almost certainly all be interpreted as secondary to the meningeal and vascular pathology. A layer of dense astrocytic glial fibres, strongly stained by the Holzer method, lay immediately beneath the pia and occupies the superficial 100–200 μm of the molecular layer. The remainder of the molecular layer showed an increased number of astrocytes and in places these were of the reactive type with abundant eosinophilic cytoplasm. Several areas of cortex showed ischaemic changes, the most severe being around the depths of some sulci with laminar necrosis, capillary and glial proliferation and abundant fat granule cells. One of these areas of cortical necrosis was much older than the others and shows abundant collagen formation within the softened zone. White matter beneath the necrotic areas of cortex showed astrocytic gliosis, some myelin loss and, in places, oedema.

Discussion

Benign lesions composed of fibrous elements, commonly termed fibrous dysplasias, desmoids, fibromas, and fibromatoses are exceedingly rare in the CNS and meninges. A review of the differential diagnosis of fibrous intracranial lesions indicates that the lesion we report is histologically unique.

Benign fibrous proliferations make up a variety of soft tissue masses and are generally classified into a group of lesions known as the fibromatoses. The fibromatoses form a spectrum of processes and are best defined as locally invasive, idiopathic, non-metastasising fibrous tumours which are of mesenchymal origin and recurrent after surgical excision. Among the fibromatoses are entities such as palmar fibromatosis, Dupuytren type fibromatosis, and congenital generalised fibromatosis. Histologically these lesions are viewed as neoplastic tissue which continues to proliferate, thereby becoming an
infiltrating local mass. Proliferation of fibroblastic elements can also be classified as reactive or inflammatory phenomena, as in the case of post-irradiation fibrosis, post subarachnoid haemorrhage fibrosis and nodular fasciitis. The histopathologic and aetologic variability of benign fibrous lesions make strict classification difficult. It appears however, that the leptomeningeal fibrosis we report has properties of both the fibromatoses and the reactive fibrous lesions.

To date there are less than 20 reports in the literature of benign CNS and meningeal fibromas or fibromatoses, and these form a variety of lesions ranging from the so-called fibroblastic meningioma to post-irradiation intracranial fibromas. Histologically these tumours were somewhat variable; however, each showed dense growth of fibrous tissue and stained densely for collagen and reticulin. Few mitoses were noted and there were no reports of metastases. It is of note that except for one isolated report of dural fibrosis presenting as infantile spasms, all other CNS fibrous lesions were discrete, well circumscribed intracranial tumours, often accompanied by a significant mass effect. In addition to hemispheric masses, an intra-cerebellar lesion and an intramedullary thoracic lesion have been observed. Fibrous tumours of bone, ossifying fibromas, are also known to occur intracranially, arising from the craniofacial bones. In contrast, the fibrous process in the present case is not tumorous in character and is confined to the leptomeninges and adjacent vessels. Macroscopically, the leptomeninges were thickened at surgery and histopathologically the lesion shows a diffuse proliferation of fibroblastic elements in the leptomeningeal plane.

Radiographically, pan-hemispheric low density was noted and the unusual feature of selective cortical enhancement extending from frontal to occipital lobes suggests that the process spread diffusely through the leptomeninges over the right hemisphere. The low density, which had been progressive since the previous scan, may reflect oedema and reaction to widespread zones of laminar cortical infarction which were found histologically. These zones of infarction are thought to be a result of perivascular fibrosis and the occlusion of vessels in the sub-arachnoid space and cortex. Arteritis was not noted histologically, but dilatation of the meningeal vessels was seen both on angiography and at surgery.

The lesion appeared to be chronic histologically and survey of the specimen showed varying degrees of density and organisation of the fibrous cellular elements. The symptom of right sided facial pain may indicate that this lesion was clinically long standing, as the pain could represent chronic trigeminal irritation or the effect of the documented engorgement of right hemispheric meningeal vessels.

The fibrootic leptomeninges are strikingly similar histologically to the soft tissue mass termed nodular fasciitis. This is a fascial mass composed of lobules of fibroblasts, capillaries, a scant chronic inflammatory cell infiltrate and areas representing granulation tissue in a myxoid stroma or collagen matrix. Nodular fasciitis has been called a "reactive" phenomenon in the soft tissues. In the case reported, although there was sub-pial gliosis, there was no clear source of meningeal irritation. Additionally, although this leptomeningeal lesion is histologically similar to nodular fasciitis, the clinical characteristics differ and here the lesion appeared to be somewhat more locally infiltrative. Although there is no evidence of direct extension into the cortex, perivascular fibrosis was noted in both grey and white matter subjacent to the thickened leptomeninges. This fibrosis seems to occupy the perivascular Virchow-
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Robin spaces. The presentation differs also, as the patient reported here had a slowly progressive 24 year history, while the reactive fibroses commonly have a much shorter clinical course.

Radiation therapy and chemotherapeutic agents have been implicated as aetiological factors causing aberrant growth of fibrous tissue. Quest et al report excision of a frontal fibrous tumour following cranial radiation therapy and CNS fibrosarcoma is well documented following irradiation. Methysergide, an ergotamine derivative used as an antimigraine agent, has been implicated as a cause of retro-peritoneal fibrosis. There is, however, no evidence in the literature of drug induced intracranial fibrosis in man. Subarachnoid haemorrhage is well known to cause fibrosis of the leptomeninges, usually basal, often leading to communicating hydrocephalus. In this case there was no history or clinical evidence of subarachnoid haemorrhage, irradiation, trauma, infection or the use of methysergide.

It is difficult to categorise formally this idiopathic leptomeningeal fibrosis. Developmentally, it is presumed that fibrous tumours of the CNS arise from mesenchymal precursors in the leptomeninges and dura or from perivascular connective tissue, and have therefore been called congenital neoplasms by Koos et al, rather than true neoplasms. A fibroma involving the dura has been described in a five month old child with congenital generalised fibromatosis. Conditions such as tuberous sclerosis and neurofibromatosis, which are associated with a proliferation of fibrous elements within the CNS, must be considered when interpreting the pathology in this case although the patient had none of the stigmata of these disorders.

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References


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