the pathophysiology of migraine, and dysfunction here may also be responsible for the impairment of consciousness.

In the present case, loss of consciousness soon after head injury very reasonably led to the diagnosis of acute extradural haemorrhage. In other cases, such features as drowsiness, confusion and hemiparesis following head injury have also led to admission to neurosurgical units. Such patients were formerly investigated by angiography, which often caused neurological deterioration, and findings suggestive of an avascular space occupying lesion (thought to be due to oedema) led in some cases to inappropriate surgical exploration.

Where coma or focal neurological dysfunction follow a mild head injury, and CT is negative, the diagnosis of trauma-induced migraine should be considered.

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References


Accepted 2 February 1989

Relapsing polynuropathies following classic Miller Fisher syndrome

Sir: The Miller Fisher syndrome is a well described form of acute inflammatory polyneuropathy, a variant of the Guillain–Barré syndrome. Initially reported by Fisher in 1956, this disease is characterised by the acute onset of ataxia, areflexia and ophthalmoplegia often preceded by a viral or other illness. Despite the alarming presentation, the disease usually has a benign outcome with a spontaneous and complete recovery within weeks. We report a case of classic Miller Fisher Syndrome followed by a bout of relapsing polynuropathy 8 weeks later.

A 21 year old female student presented with complaints of facial paraesthesias, diplopia, generalised weakness, and a gait disturbance described as a “loss of balance sense.”

Physical examination of heart, lung and abdomen was normal. Neurological examination revealed an ophthalmoplegia involving all extraocular muscles with minimal vertical gaze preserved bilaterally. Bilateral ptosis was noted, with no nystagmus. Pupils were reactive to light and accommodation. Deep tendon reflexes were slight in the upper extremities and absent in the lower extremities. Her upper and lower extremity strength was intact bilaterally, as was sensory perception. Ataxia was present, with a wide based, unsteady gait.

Lumbar puncture revealed a glucose of 58 mg/100 ml, protein of 72 mg/100 ml and a cell count of 10/mm3 RBC and 1/mm3 WBC. Nerve conduction and EMG studies were performed. Skin temperature was within the normal range (31°C). Peroneal nerve motor velocity was 44 m/s. F wave conduction velocities were slow at 35 mm/s. Sural responses were absent despite averaging. This was consistent with a mild demyelinating neuropathy.

The patient was diagnosed as suffering from Miller Fisher syndrome and was observed in the intensive care unit. Over the next several days, the patient experienced progressive leg and arm paraesthesias, without a deficit in limb strength. Slowly, the patient improved, and was discharged 10 days after admission with minimal ptosis, paraesthesias, facial weakness and ataxia.

The patient was seen one month later where she was found to be walking easily and feeling stronger. There was full ocular motility, with minimal ptosis of the left eye and mild right sided facial weakness. Sensory testing revealed minimally diminished vibration and position sense of hands and fingers with mildly diminished pin sensation in a stocking distribution. The gait was normal. Deep tendon reflexes were absent.

One week later she developed an exacerbation of her neurological symptoms. Ambulatory skills declined with development of foot and hand paraesthesias. On re-examination the patient was noted to have no position sense with manual testing of her hands and feet or any distal vibratory sense. Strength was good and her extraocular muscles remained intact. She had no exacerbation of her facial weakness or ptosis. Her dramatic clinical decline was consistent with a disabling relapsing polyneuropathy, primarily sensory in nature. She was then started on a two week regimen of oral prednisolone.

After one week of steroid therapy, the patient reported a dramatic improvement in her symptomatology. Unsteadiness, imbalance and diminished position were less noticeable. It was felt that she was entering a remission stage of her neuropathy.

Physical examination three months after her initial diagnosis revealed persistence of the absent deep tendon reflexes. The facial weakness and ptosis, significant at the onset of her illness, were no longer apparent. Extraocular muscle movement was without limitation. Tactile sensation was normal, although position and vibration were still mildly impaired. She was able to walk without assistance and anticipated returning to her activities of normal living.

The Miller Fisher syndrome is a rare form of acquired demyelinating neuropathy. A variant of Guillain–Barré syndrome, this monophasic illness develops acutely over a few days. The ataxia is often severe and debilitating. Symmetrical ocular palsies occur with greater involvement of external musculature. Limb involvement is minimal or absent, although motor weakness has been known to occur. Patients usually recover without specific treatment over the course of several weeks after a relatively benign but frightening illness.

Our patient exhibited all the features of the syndrome. What distinguished our case from the others was the development of recurring polynuropathies complicating the initial diagnosis of Miller Fisher syndrome. We did not expect her to relapse after her rather benign initial course. Interestingly, during the time of the relapse, the patient did not exhibit any exacerbation of her ophthalmoplegia or facial weakness. Her main problem was sensory with loss of position and vibratory sense of her extremities, as well as pin touch. A dramatic improvement was noted after the introduction of corticosteroid therapy.

A review of the literature has yielded similar but not identical cases. Donaghy et al reported three cases of oculomotor palsy preceding the development of chronic polyneuropathies by an interval of two to ten weeks. Two patients with ophthalmoparesis and recurrent sensory neuropathy were reported by Kaplan. Chalmers and Miller in 1986 reported a patient with a 2 year history of chronic progressive inflammatory polyneuropathy with a subacute exacerbation that included ataxia, areflexia, ophthalmoplegia and ptosis. Schapira reported a patient with a relapsing neurological deficit. He initially manifested the Miller Fisher syndrome after a viral illness.
syndrome with Guillain-Barré features (limb weakness and distal paresthesias), and during a relapse one year later, exhibited purely Miller Fisher features.

Our patient had a classic manifestation of Miller Fisher syndrome with ataxia, areflexia and ophthalmoplegia followed by a recurring polyneuropathy eight weeks later. We believe this is the first reported case of this association. Most often the course of Miller Fisher syndrome is one of spontaneous recovery. This patient reinforces the need for continued observation of these patients, even after initial improvement of their neurological deficit.

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References

Spontaneous haemorrhage in craniopharyngioma

Sir: While chemical meningitis caused by the extravasation of oily cystic fluid from a ruptured craniopharyngioma is known, it usually occurs after surgery. Spontaneous rupture is more rare and proven haemorrhage has not been documented in this tumour. We describe a patient who presented with subarachnoid haemorrhage secondary to haemorrhage in a craniopharyngioma.

A 59 year old housewife, who had been suffering from sciatric pain and mild headache for two months developed severe headache and nausea on the day before admission. She had no contributory previous medical or family history. On examination the patient was alert, cooperative but was in some distress. The detected abnormalities were mild fever and pain bilaterally upon straight leg raising. Her neck was supple.

Lumbar puncture yielded deeply xanthochromic, turbid cerebrospinal fluid (CSF) with the opening pressure of 22 cm H2O. Other than the uncounted erythrocytes, the cell number of the CSF was 250 per mm3 with 59% polymorphonuclear leukocytes, 38% lymphocytes, and 3% mononuclear cells; the protein was 125 mg/dl, and the glucose 38 mg/dl. There were no detectable cholesterol crystals. The CSF cytology was negative. CT revealed a high density suprasellar mass (fig A) which was diffusely enhanced by contrast medium. Plain radiographs of the skull demonstrated calcification in the suprasellar region corresponding to the mass in CT. The cerebral angiogram was unremarkable. The tentative diagnosis was a haemorrhage either into a craniopharyngioma or a pituitary adenoma. Laboratory examination of the peripheral blood was unremarkable except for the mildly elevated prolactin level.

Repeat CT without contrast medium revealed a gradual decrease in the density of the juxtasellar portion of the mass (fig B).

Two months after admission, subtotal removal of the suprasellar tumour was performed via the right fronto-temporal approach. The arachnoid membrane was brownish and thick. Behind the optic chiasm was a partly dark coloured, partly calcified mass which did not yield any fluid. Histological examination of the specimen demonstrated typical craniopharyngioma consisting of islands of epidermoid tissue in the adamanitomatosum pattern, squamous cells, keratinisation, calcification, and microcyst formation. In addition, there were blood clots and numerous haemosiderin-laden macrophages. Some of the blood vessel walls in the tumour and the surrounding connective tissue stroma showed hyaline or fibrous thickening. The postoperative course was uneventful, except for transient diabetes insipidus.

The clinical manifestations of craniopharyngioma include visual field deficit, hypopituitarism, increased intracranial pressure caused by compression of the surrounding structures and chemical meningitis due to extravasation of cystic fluid. However, haemorrhage has not been confidently proved to occur in craniopharyngioma as it has, rarely, in pituitary adenoma.

As in the present case, the CT density of some of craniopharyngiomas has been reported to vary with time. In such a case the CT change was attributed to the protein concentration, not to haemorrhage, in the cyst. In another case haemosiderin deposition was found in the connective tissue stroma of the cyst wall. The authors suggested haemorrhage in the tumour although there had been no sudden episode of clinical deterioration or bloody cyst fluid at surgery. The CSF and preoperative CT findings of

Fig. CT without contrast medium, showing a juxtasellar high density mass (A) 1 day and (B) 42 days after the onset of symptom.
Relapsing polyneuritis following classic Miller Fisher syndrome.

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*J Neurol Neurosurg Psychiatry* 1989 52: 802-803
doi: 10.1136/jnnp.52.6.802

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