Posterior ischaemic optic neuropathy after a spontaneous extradural haematoma

Most extradural haematomas (EDH) occur after head injury but cases of “spontaneous” EDH have been reported. These are a result of either local sepsis, a dural arteriovenous malformation or an abnormality of clotting. We report the case of a spontaneous EDH resulting in posterior ischaemic optic neuropathy as a consequence of tentorial herniation.

A 15 year old boy developed a headache which built up over a period of 30 minutes. In the evening he was seen by his general practitioner who diagnosed a viral infection as the headache was associated with some fever. On the second day he vomited in the morning. By the morning of the third day he had fever and was unable to get up because of headache and giddiness. His mother saw him at approximately 1.00 am on the fourth day of his illness when he seemed to be sleeping normally. By 7.00 am the same morning he was found unconscious and was admitted to the District General Hospital on 17 March 1988. There was no other predisposing illness or head injury.

In the accident and emergency department, he was noted to be cyanosed with vomit in his nose and throat. He was in coma with a Glasgow Coma Score of 5. Both pupils did not respond to light and the right was 6 mm, the left 3 mm. His axillary temperature was 40°C, a regular pulse of 110, blood pressure 120/55 mmHg and normal heart sounds. Auscultation of his chest revealed added sounds in the left base suggesting an aspiration pneumonia which was confirmed on chest x ray. His blood results were as follows: HB 1·38 g/l, white cell count 11·7 × 10⁹/l, platelet count 180 × 10⁹/l, sodium 140 mmol/l, potassium 4·2 mmol/l and urea 4·1 mmol/l. Clotting was within normal limits with an INR 1·0 and KCCT (control 48).

Before being transferred, the patient was intubated, ventilated, paralysed, sedated and given 100 ml mannitol 20%. On arrival in the neurosurgical unit on 17 March 1988, a CT was performed (figure) which showed a 3 cm maximum depth EDH over the right frontal convexity which was of mixed density. There was no radiological evidence of head injury. In view of his condition and the radiological findings, an immediate right frontal craniotomy was performed and a moderately sized EDH was found. The haematoma was considered to be of some days duration. No evidence was found of either a fracture or a dural arteriovenous malformation.

On the evening of admission the patient developed a transient diuresis which resolved and there were no further episodes. The patient was extubated four days postoperatively and his level continued to improve. Initially his vision was very restricted and he could see only a few objects. Examination showed that the pupils, although unequal, were reacting to light. He appeared to have a left homonymous hemianopia very early after surgery. This was not demonstrable subsequently as he has remained completely blind since then. He was also noted to have a mild left hemiparesis (Medical Research Council grade 4). One month later he was discharged and his conscious level had returned to normal, his hemiparesis had resolved but he was blind.

On review to the accident department three months later, his vision was unchanged with no light perception in either eye. He had put on 19 kg in weight. He had polydipsia and, according to his mother, prolonged and excessive periods of sleeping. On examination he was alert and orientated with no limb weakness. His pupils did not react to light and funduscopy revealed bilateral optic atrophy. Visual evoked responses showed no response. A CT scan showed moderate atrophy of the right frontal lobe and mild dilatation of the ventricular system compared with the previous scan. The orbits, sella and suprasellar cisterns appeared normal. MRI showed slender optic nerves consistent with atrophy and confirmed the frontal atrophy. There was no evidence of hypothalamic abnormality on the scan. He had an endocrine assessment six months postoperatively which showed an isolated defect of growth hormone production.

One year postoperatively the patient remains blind with fixed pupils protect subsistent signs from MPP—neurotoxicity in rats. Injection of the NMDA antagonist MK 801 within the medial pallidum reverses Parkinsonian symptoms in MPTP-treated monkeys. The mechanism of action remains unclear but it has been suggested that blockade of NMDA receptors would result in a facilitation of dopamine action by preventing the glutamate-induced dephosphorylation of DARPP-32, a dopamine- and cAMP-regulated phosphoprotein. However, no clinical data are yet available in Parkinson’s disease (PD).

We investigated in PD patients the effects of an add-on therapy with ifenprodil, a non competitive antagonist of the NMDA receptor at the polyamine modulatory site, which inhibits [3H] MK 801 binding. This drug is, as far as we know, the sole NMDA antagonist currently available on the market. Ifenprodil also possesses alpha-adrenoceptor blocking properties and is already in clinical use as a central antihypertensive agent.

Two groups of patients with idiopathic PD were studied after informed consent: the first one included nine non-fluctuating patients, mean age (SD), 67 (14) years, and SD duration of PD 9 (1) years, mean (SD) duration of levodopa therapy: 7 (1) years,

**Figure** CT scan showing a 3 cm extradural haematoma of mixed density in the right frontal region.
meas (SD) dose of levodopa : 595 (88) mg, mean (SD) stage of Hoehn and Yahr : 3-2 (0-3) with waning efficacy of levodopa and who were requiring an increase in PD treatment. The second group included 11 other patients, mean age, 61 (2) years, mean (SD) duration of PD : 12 (1) years, mean (SD) dose of levodopa : 857 (92) mg, mean (SD) duration of levodopa therapy : 10 (1) years, mean (SD) stage of Hoehn and Yahr : 3-0 (0-3) with peak-dose dyskinesias most of the day. In both groups, ifenprodil (Vadilax), (20 mg three times daily orally, the dose currently used in clinical practice) was added in an open design without changing any previous treatments, usually during PD treatment. These two kinds of PD patients were selected to investigate if the NMDA antagonist could improve Parkinsonian symptoms (group 1) or modify dyskinesias (group 2). Parkinsonian symptoms were evaluated in the morning (10 am) according to the Unified Parkinson's Disease Rating Scale (UPDRS) before and after 1 month of treatment. Each new assessment was made blind, without the previous scale score in front of the physician. Patients of group 2 were assessed in the “on” condition. Add-on therapy with ifenprodil did not modify the Parkinsonian symptoms in any group. The total UPDRS motor examination score did not change significantly in group 1 186 (3-5) vs 20-2 (3-2) or group 2 29-4 (2-3) vs 8-6 (2-6). The UPDRS subscores for cardinal extrapyramidal symptoms (tremor, rigidity, bradykinesia) or for daily activities did not change (data not shown). The dyskinesia score remained unchanged in group 2. Side effects were palpitations and sedation (1 patient) and feeling of nasal congestion (1 patient).

Our study is the first to investigate the clinical effects of an NMDA antagonist in the treatment of PD. It failed to demonstrate any relevant anti-Parkinsonian effect of ifenprodil. These negative results must, however, be considered with caution. We used the daily dose of ifenprodil recommended for the treatment of intermittent claudication but the pharmacokinetic profile of the drug is poorly known. No published data are available about its plasma half life and brain distribution, making it possible that other NMDA antagonists with a better pharmacokinetic profile may exert more potent effects in PD. Since the polyamine modulatory site is only a part of the NMDA receptor complex, it is also difficult to give a precise biochemical interpretation of our result. Our work does not exclude a definite role for NMDA antagonists in PD. Further studies should be conducted with other NMDA antagonists when available in humans.

JL MONTASTRUC, O RASCAL, L MAURON, A RASCAL*
Department of Neurology, hen Hospital Universitaire Purpan, faculté de Médecine, 37 allée Jules-Guesde, 31073 Toulouse, France.


Central pontine myelinolysis in a patient with AIDS

Central pontine myelinolysis (CPM) has been associated with rapid correction of hyponatraemia as well as an acute rise in serum sodium even from normonatraemic levels.4 CPM has also been linked with hyperosmolality,5 the rate of change of osmolality, the length of time that this rate of change persists, and the clinical condition of the patient are all important factors.6 We describe a case of CPM in a 49 year old HIV positive homosexual man who had a normal serum sodium and osmolality throughout his illness.

The patient had been HIV positive for at least six years. In 1989 he developed increasing hepatoosmologenomally uncontrolled thrombocytopenia. A splenectomy was performed and histology showed only non-specific changes associated with HIV. After an initial stormy postoperative course, he made an excellent recovery and returned to work.

Seven months later he presented with a short history of fevers, lethargy, and fever.

The only abnormal finding was a pericardial rub. Investigations showed that he was mildly anaemic, and hyypoalbuminaemic with a serum albumin of 19 g/l. Liver function tests and clotting studies were normal. Initial investigations were all negative apart from demonstrating a pericardial effusion and a small pleural effusion. On rescreening for pericardiocentesis, however, the effusion had resolved. A small pleural effusion was aspirated and found to contain numerous lymphocytes suggesting a lymphoma. Immunocytochemistry was unhelpful. Bone marrow examination and 24 hour urine protein excretion were normal. He had no gastrointestinal symptoms. Thirteen days before his death he developed diplopia and was found to have a right sixth nerve palsy. A CT brain scan showed no abnormality. A lumbar puncture showed clear CSF with 10 lymphocytes and a protein of 0-78 g/l. No pathogens were isolated and he was cryptococcal antigen negative. Cytological examination of the CSF showed no abnormality. A trial of high dose steroids made no difference to his deterioration. He became progressively weaker, developed right-sided posis and dysarthria and died a few days later.

Necropsy examination demonstrated disseminated lymphoma, including large ulcerating deposits in the stomach. Postmortem examination of the brain showed the typical appearance of CPM with a large well defined focus of myelin loss in the base of the mid-pons and upper-pons (figure). Focal infiltrates of lipid containing macrophages were present in the area of myelinolysis but there was no inflammation otherwise associated with this lesion. Axons were partly preserved. There was some mild cerebral atrophy, mild diffuse myelin pallor in the cerebrum and slight lymphomatous infiltration of the meninges.

To our knowledge there have been no previous reports of CPM in association with HIV. The only significant biochemical abnormality was the hypoalbuminaemia which developed in the month preceding his death. This may have been due to gastro-intestinal losses as well as to his underlying condition. His serum sodium remained normal throughout his stay in hospital, and he received no intravenous fluids. His albumin remained at 19 g/l until he died. CPM has been described in liver transplant recipients and in severe burns cases, as well as in association with alcoholism and malnutrition. It has been thought that the rapid correction, or over correction, of hyponatraemia was the crucial factor in causing CPM in most cases. This is obviously not the only cause. In a series of CPM in severely burnt patients, hyponatraemia was not a feature, whereas each patient who developed CPM had a prolonged episode of extreme hyperosmolality. Our patient did not have an episode of serum acute hyperosmolality, but had one month of hyperosmolarinaemia. A low serum albumin may be a significant factor in the development of CPM. This could be a feature common to our patient and the groups of patients described above in which CPM had been reported. Another possibility is that CPM may be a primary effect of HIV, representing yet another neuropathological manifestation of this virus. The p24 HIV protein was detected immunocytochemically in rare perivascular macrophages.