LETTERS TO THE EDITOR

Ophthalmoplegic migraine and recurrent sinus arrest

Cardioinhibitory vagovagal reflex, induced by pain, is well known. We are unaware, however, from the literature, of recurrent sinus arrest episodes in the setting of ophthalmoplegic migraine, a rare entity in itself (Kandt et al. 1960).

A 34 year old female was admitted to the intensive cardiac care unit on 19 July 1987 following three events of dizzy spells, malaise and near fainting. The day before she sustained the right of a severe headache, either unilateral or bilateral, sometimes accompanied by nausea. She did not smoke or take any medication. We found no family history of headache, seizure, sudden death or chronic diseases.

During six consecutive days in the intensive care unit, she sustained recurrent pain in her left temple and almost every time a sinus arrest was recorded on the monitor, 10-20 times per day.

On the sixth day, the headache began to recede and gradually left sixth nerve palsy and diplopia became apparent as the only neurologic deficit. EEG and brain CT, as well as LP, performed when a sixth nerve palsy appeared, were normal. Her double vision completely cleared three weeks later.

A trial of atropine treatment (iv) was given, resulting in a switch from extreme sinus bradycardia to regular nodal rhythm. The patient refused invasive electrophysiological study of the heart. Autonomic functions were normal: her blood pressure ranged between 110/70 and 120/70 mmHg. Orthostatic hypotension was absent. Her diaphoresis was not disturbed and bowel movement was regular. Limited tests were done. Eyeball reflex did not change the heart rhythm in the interictal periods; carotid massage did not reveal hypersensitivity of the carotid sinus. Valsalva manoeuvre showed a normal “overshoot” of blood pressure in the fourth phase.

The diagnosis was based on the clinical course, outcome, physical examination, normal CSF and brain CT. On CT with enhancement, no aneurysm was visualised. The patient refused carotid angiography.

Tolosa-Hunt painful ophthalmoplegia is another possibility. Hunt’s criteria were almost totally fulfilled. It is, however, difficult to differentiate between the two conditions. The short duration of pain and the benign self-limiting course favoured the diagnosis of ophthalmoplegic migraine. The speculative explanation of the mechanism in the two entities indicates that they may share common features.

Symptoms of autonomic dysfunction are well known in migraine, and an ophthalmoplegic variant is not an exception in this case. This autonomic dysfunction was an isolated episode. Abnormalities in the sympathetic nervous system in those with migraine headache are well known, but cause arthralgias, and not sinus arrest. The right vagus is supposed to innervate the sinoatrial node; its stimulation causes sinus arrest. It is not clear how it was stimulated by a presumed left extra-axial process. Perhaps the pain acts as “non-localised” trigger of multiple sinus arrest spells.

A follow up over two years found the patient healthy, except for her “habitual” headache, and no focal signs have been detected.

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Reversal of foot drop in sciatic nerve endometriosis

Pelvic endometriosis is a common gynaecological disorder. While usually involving the ovaries, uterosacral ligaments and pelvic peritoneum, it may rarely invade the sciatic nerve and its plexus. The latter complication is typified by cyclical sciatica or foot drop. Previous endeavours to treat leg weakness in pelvic endometriosis have met with limited success. In our study of sciatic nerve endometriosis we report the successful reversal of foot drop with danazol.

A 46 year old single nulliparous woman presented to the neurology service with a twenty month history of progressive, painless right foot drop. For thirteen months she had experienced numbness in the right foot and for several weeks tingling in the right sole. She gave a history of a severe head injury at age 12 years but had recovered well with only mild balance problems thereafter.

The patient had a long-standing history of regular heavy menses associated with troublesome dysmenorrhoea.

On examination there was severe weakness of dorsiflexion and plantar flexion of the right foot and toes and moderate weakness of the hamstrings. The right ankle reflex was absent and the right foot cold and blue. Tone was reduced at the right ankle but there was no muscle wasting and no fasciculations. Light touch, pain and temperature were reduced over the dorsum and plantar surface of the right foot extending laterally to the mid calf level. Vibration and position sense were intact. Electromyographic examination of the right tibialis anterior, peroneus longus, medial hamstring and glutaeus medius muscles showed evidence of active denervation with fibrillation potentials, increased numbers of polyphasic motor unit potentials and impaired motor unit recruitment. No evidence of denervation was found in the right vastus lateralis, glutaeus maximus and lumbar sacral paraspinal muscles. A myelogram was normal but a pelvic ultrasound scan revealed a large complex intrapelvic mass measuring 11 × 9 × 9 cm which appeared malignant.

At laparotomy, severe endometriosis was revealed with an American Fertility Society Score of 136/150. There was complete obliteration of the pouch of Douglas with a large left ovarian endometrioma and a smaller fixed right tubo-ovarian endometriotic mass. A left salpingo-oophorectomy was performed but due to fixation on the right, the endometrioma was drained but not excised.

A post operative pelvic CT scan revealed a 5 × 3 cm hypodense mass adjacent to the right sacro-iliac joint (fig 1). The patient was started on danazol 200 mg twice daily.

Three months after surgery she no longer had paraesthesia and dorsiflexion of the right foot was only slightly impaired. At twelve months, dorsiflexion of the right foot was normal and the right ankle reflex had returned. Light touch, pain and temperature were impaired only distal to the right ankle. Electromyographic studies showed evidence of recovery with an absence of fibrillation potentials in previously denervated muscles in the right leg. Furthermore, these muscles exhibited increased amplitudes and increased “jitter” indicative of active collateral sprouting. Danazol treatment was stopped after eleven months.

A CT pelvic scan at six and 12 months showed no significant change in American Fertility Score of 136/150. There was complete obliteration of the pouch of Douglas with a large left ovarian endometrioma and a smaller fixed right tubo-ovarian endometriotic mass. A left salpingo-oophorectomy was performed but due to fixation on the right, the endometrioma was drained but not excised.

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