Letters

Conjunctive deviation of gaze in hepatic encephalopathy

Sir: Transient ocular motor disturbances such as ocular bobbing,1 dysconjugate gaze,2 skew deviation3 and absent horizontal responsiveness to oculovestibular testing4 have been described in patients with hepatic encephalopathy. Sustained conjunctive deviation of the eyes, however, is usually considered indicative of a contralateral irritative lesion or an ipsilateral destructive lesion of the cortex or a lesion of the contralateral pontine gaze centre.5 Here we describe a patient with fulminating hepatic failure in whom conjunctive deviation of gaze developed two days before death. No structural lesion could be demonstrated on CT scan or at necropsy.

A 52-year-old man with a long history of ethanol abuse and biopsy-proven hepatic cirrhosis was admitted in an obtunded state. On examination he had tense ascites and pedal oedema and heavily jaundiced sclerae. He responded to only simple verbal commands and had bilateral asterixis; neurological assessment was otherwise unremarkable. Laboratory data at that time were: total bilirubin 273 mmol/l; serum albumin 16 g/l; serum aspartate transaminase 518 U/l, lactate dehydrogenase 452 U/l; alkaline phosphatase 915 U/l and y-glutamyltransferase 430 U/l; haemoglobin 9.3 g/dl. Treatment with low protein diet, lactulose, and spironolactone was instituted. The patient’s mental state deteriorated progressively over the next week and he became unresponsive to verbal stimuli but continued to respond purposefully to pain. At this time it was noticed that his eyes were deviated to the left in a conjunctive fashion and failed to respond to caloric and oculovestibular stimuli. Pupils were equal and reactive to light and there was no papilloedema. Plantar responses were flexor and tone of the limbs was normal. Total bilirubin was now 316 μmol/l and serum ammonia 116 μmol/l (normal 17-47 μmol/l).

A structural cortical or brainstem lesion was suspected. CT scan was normal and EEG showed nonspecific slowing in the theta range. CSF analysis showed only the presence of bilirubin in the fluid. The eyes remained deviated until two days later when the patient died suddenly. The last total bilirubin level a few hours before death was 522 μmol/l.

A necropsy was performed. The brain (wet weight 1380 g) was jaundiced, but the most careful macroscopic and microscopic examination failed to reveal any structural lesion in any area. In keeping with the clinical picture, the liver was shrunken (840 g) and cirrhotic.

Stupor in this patient was due to hepatic encephalopathy. Cirrhosis with ascites proved by biopsy and necropsy, pedal oedema, abnormal liver function tests with elevated serum ammonia, diffuse EEG slowing and normal CT scan and CSF make this conclusion inescapable. The presence of sustained conjunctive deviation of gaze, not previously described in hepatic stupor, was confusing and suggested a structural lesion. No such lesion could be demonstrated, either on CT scan or at necropsy. The occurrence of the conjunctive deviation in this case must, presumably, be explicable on the basis of selective vulnerability to metabolic insult of centres associated with control of conjunctive eye movements.

There is no doubt that toxic-metabolic conditions may affect brain-stem tegmental function as well as motor pathways and hemispheric structures.2 3 In such patients, systemic and laboratory evidence of severe metabolic disturbance and preservation of other functions at the same level should suggest a metabolic explanation for the signs.

R SANDYK
MIW BRENNAN
MMW ERDMANN
Department of Medicine,
Hillbrow Hospital,
Johannesburg 2001,
South Africa

References


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Recurrent meningitis due to labyrinthine fistula

Sir: We wish to describe two children who had recurrent meningitis. In children recurrent meningitis may have various causes.1-3 In the absence of an immunological defect, it may be the result of anatomical abnormalities, such as skull fracture, tumour and congenital defects. In a few cases, the pathology responsible for the recurrent meningitis may be in the middle ear.4-6 The triad of recurrent meningitis, severe unilateral or bilateral sensori-neural deafness and opaque tympanic membrane or other evidence of fluid in the middle ear should raise the possibility of middle ear pathology. The two children we describe, both presented with this triad arising from labyrinthine fistula.

A 5-year-old girl with a normal birth history and development presented initially at 7 months of age with Haemophilus influenzae meningitis. She recovered quickly but developed severe bilateral sensori-neural deafness. A few weeks later, she had an attack of pneumococcal meningitis which recurred at the age of 23 months. Radiological examination revealed a defect at the apex of the petrous temporal bone. A left lateral craniotomy with dural repair6 of the middle fossa was performed at 2 years of age. However, she had five further attacks of meningitis which included three due to pneumococcus and two due to Haemophilus influenzae between 2 years 3 months and 6 years 6 months. At the age of 5 years 8 months she was admitted to the Hospital for Sick Children, London. Examination showed the left tympanic membrane to be normal, but the right was bluish and a fluid level was seen. She had bilateral sensori-neural deafness. The rest of the examination, including full neurological examination, was normal. The results of investigations were not diagnostic: haemoglobin level was 11 · 9 g/dl; white cell count was 8.7 × 109/l (neutrophils 71%; lymphocytes 21%; reticulocytes 5%); eosinophils 2%; basophils 1%).

Complement C3 was 108% of standard. Nitroblue tetrazolium test was normal. Radiographs of the sinuses showed minor mucosal thickening in the antrum. Chest radiograph was normal. Tomograms of the mastoids showed the lateral part of the right internal auditory meatus wider than that on the left, and the vestibule was slightly larger. The basal coil of the right cochlea was normal. Computed tomography (CT) showed an opaque right middle ear. A right exploratory