Absent vestibulo-ocular reflexes and acute supratentorial lesions

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SYNOPSIS Loss of vestibulo-ocular reflexes occurred in two patients with acute supratentorial lesions who received therapeutic doses of anticonvulsant drugs. There was no clinical or angiographic evidence of focal brain-stem damage. Absence of vestibulo-ocular reflexes is attributed to a combination of acute cerebral damage and anticonvulsant drugs. The loss of these reflexes in patients with acute cerebral lesions cannot be interpreted as evidence of irreversible brain-stem injury.

Loss of vestibulo-ocular reflexes may result from structural brain-stem lesions, drug intoxication, metabolic encephalopathy or peripheral vestibular disease (Rodríguez Barrios et al., 1966; Plum and Posner, 1972). Acute lesions of the cerebral hemispheres may extinguish the fast phase of induced vestibular nystagmus (Merwan and Feiring, 1939), but loss of all vestibulo-ocular response has been considered to be an indicator of advanced brain-stem damage secondary to transtentorial herniation (Vaernet, 1957). This paper reports reversible loss of vestibulo-ocular reflexes with acute supratentorial lesions in two patients receiving therapeutic doses of anticonvulsant drugs.

CASE 1

An 11 year old boy, with a previous diagnosis of right middle cerebral arteriovenous malformation, suddenly developed severe headache, then flaccid left hemiplegia and stupor. He had one seizure, for which he received an intramuscular and intravenous injection of diphenylhydantoin (Dilantin) and intravenous phenobarbitone at an emergency hospital. Two days later, still stuporous and hemiplegic, he was transferred to the neurosurgical service of the University of California, San Francisco (UCSF).

On command, the boy made purposeful withdrawal movements of the right arm and on occasion made appropriate verbal responses to questions. His respiratory pattern was normal. Blood pressure was 115/80 mmHg, pulse rate 82 per minute, and rectal temperature 38°C. Left hemiparesis persisted. He had paraparesis of the right arm and leg. Although his eyes were closed, bilateral blink reflexes were elicited by touching the margins of the eyelashes. The patient's corneal reflexes were normal. Pupils were 4 mm in diameter and reacted briskly to light. His eyes were in mid position and did not move spontaneously. Ophthalmoscopic examination showed optic disc and nerve fibre patterns of homonymous hemiopic hypoplasia. Spontaneous venous pulsations were absent, but there was no disc oedema.

NEURORADIOLOGICAL FINDINGS The internal cerebral vein was 3 mm to the left of midline in the left internal carotid angiogram. No mass effect was present in the lateral projection of the right carotid angiogram. The arteriovenous anomaly was 2 × 3 cm in the lateral projection, being situated deep in the parietal lobe and extending along the wall of the lateral ventricle. There were no angiographic signs of transtentorial or tonsillar herniation, and there was no rostral caudal shift of the brain-stem. There were no angiographic indications of acute obstructive hydrocephalus.

VESTIBULO-OCULAR FINDINGS Rapid turning of the head to either side evoked no oculocephalic res-
responses. Passive movement of the head upward and downward in the vertical plane stimulated conjugate vertical eye movements 10° both below and above the horizontal meridian. Ice-water irrigation of each external auditory canal combined with rapid head turning did not produce eye movement horizontally in either direction. Examination of the external auditory canals was normal. Repeat caloric tests were unchanged in three hours. Serum determinations of diphenylhydantoin and phenobarbitone obtained at this time were 2.5 mg and 2.9 mg per 100 ml, respectively.

On the patient’s third day in the hospital, oculocephalic (‘doll’s head’) testing produced eye movements 5° to the right, 5° to the left, 20° up, and 20° down. Ice-water caloric testing produced the same horizontal ocular deviations. Serum levels of diphenylhydantoin and phenobarbitone were unchanged. By the fourth day, the patient was more alert, though his verbal communication remained meagre. Oculocephalic and caloric tests now produced ocular tonic deviation all the way to the right, but still only 5° to the left; eye movements in the vertical direction were unchanged. Serum levels of diphenylhydantoin and phenobarbitone were 2.7 mg and 2.5 mg per 100 ml.

**TREATMENT AND COURSE.** While he was in the hospital, the patient received diphenylhydantoin (100 mg three times daily), orally. Phenobarbitone (30 mg four times daily) was administered orally until resection of the arteriovenous malformation on the tenth hospital day. During surgery old clotted blood was seen in the right ventricle.

After operation the patient voluntarily moved his eyes completely to the right, but he could not move them beyond the midline to the left. No leftward ocular movement could be elicited by turning his head. There was no vertical movement of the eyes with upward and downward movement of the head.

Nine days after surgery, the patient voluntarily moved his eyes 30° up, 30° down, 30° to 40° to the right, and 25° to 30° to the left; two weeks after operation he voluntarily moved his eyes fully in all directions. The patient was discharged five weeks after admission with persisting left hemiplegia and hemianopia.

**CASE 2**

A 22 year old man had a febrile `flu-like’ illness followed two weeks later by headache, nausea, vomiting, and five serial grand mal seizures. He was admitted to a local hospital and diphenylhydantoin (200 mg twice daily) and phenobarbitone (30 mg four times daily) were given by nasogastric tube for frequent seizures. The patient moved his arms and legs in response to painful stimuli. His pupils were equal and reacted briskly to light; his optic discs were normal and his eyes moved conjugately and completely in vertical and horizontal directions. Occasionally he had tonic conjugate movements of his eyes to the right, for one to two minutes. Lumbar puncture showed a mildly xanthochromic cerebrospinal fluid at a pressure of 210 mm H₂O with one red blood cell, 11 white blood cells/ml (four polymorphonuclear leucocytes, seven lymphocytes), glucose 31 mg and protein 38 mg per 100 ml. An electroencephalogram showed epileptiform discharge over both temporal areas. Seizure activity persisted, and ethosuximide (Zarontin) (200 mg three times daily) was given by nasogastric tube. Serum diphenylhydantoin and serum phenobarbitone measured two days after admission were 2.3 mg and 0.9 mg per 100 ml respectively. Serum electrolytes at this time were Na 147 mEq/l, K 3 mEq/l, Cl 102 mEq/l, and HCO₃ 24 mEq/l. Seizures continued and he was treated with glycerin (90 oz three to four times daily) by nasogastric tube, hydrocortisone (250 mg four times daily), sodium amylobarbitone (25 mg three to four times daily) and diazepam (5 mg three to four times daily) intravenously. Four days after admission serum electrolytes were Na 163 mEq/l, K 2.9 mEq/l, Cl 120 mEq/l, HCO₃ 32 mEq/l, and the blood urea nitrogen was 74 mg per 100 ml. Diphenylhydantoin and ethosuximide were discontinued and he was given intravenous fluid replacement. He was transferred to the neurology service of UCSF Hospital on the fifth day after admission; during the preceding 24 hours, the patient had received diazepam (35 mg) and phenobarbitone (200 mg) intravenously.

At initial examination, he was unresponsive to verbal stimuli but made defensive withdrawal movements of both arms and legs in response to painful stimuli. Generalized seizures occurred almost hourly. Respiration were normal. Blood pressure was 150/70 mmHg, pulse rate 102/min, and rectal temperature 38.5°C. His limbs were flaccid. Reflexes were symmetrically brisk, and plantar responses were absent. The patient’s eyes were closed but bilateral blink reflexes were elicited by touching his eyelashes. His corneal reflexes were symmetrically depressed. The right pupil was 5 mm, the left pupil 4 mm in diameter, and they reacted briskly to light. His eyes were in mid position and did not move. Optic discs were flat and spontaneous venous pulsations were present.

**NEURORADIOLOGICAL FINDINGS.** Bilateral carotid angiography showed no filling of the superior or inferior sagittal sinuses. There was prolonged
opacification of convexity veins. The posterior venous circulation was normal. No midline shift was evident and the ventricles were of normal size.

VESTIBULO-OCULAR FINDINGS Rapid turning of the head to either side and upward and downward did not elicit any eye movement. Combined ice-water irrigation of each external auditory canal and head turning also failed to elicit eye movement. The external auditory canals were not obstructed. Serum determinations of diphenylhydantoin and phenobarbitone at this time were 2.5 mg and 2.0 mg per 100 ml respectively.

On the third day of hospitalization at UCSF, the patient voluntarily moved his eyes about 15° to the right and 15° to the left. These movements were slow and there was gaze-evoked nystagmus in both directions. He could not move his eyes up or down. At this time the serum level of diphenylhydantoin was 2.5 mg/100 ml and serum phenobarbitone was 3.0 mg/100 ml. However, by the fifth day, voluntary movements were full, with gaze-evoked nystagmus in all directions. Determinations of serum diphenylhydantoin and phenobarbitone by this time were 1.0 mg and 5.9 mg/100 ml.

TREATMENT AND COURSE. Sodium amylobarbitone (170 mg), diazepam (35 mg), and phenobarbitone (400 mg) were given intravenously on the day of admission and the seizures were controlled; thereafter diphenylhydantoin (400 mg daily) and phenobarbitone (150 mg daily) were given to maintain control. A lumbar puncture showed an opening pressure of 420 mm H₂O and closing pressure of 210 mm H₂O after removal of 6 ml of cerebrospinal fluid containing protein 163 mg/100 ml, glucose 78 mg/100 ml, 30 red blood cells/ml, and no white blood cells. Serum electrolyte determinations were Na 138 mEq/l, K 3.9 mEq/l, Cl 106 mEq/l, HCO₃ 28 mEq/l, Ca 8.6 mg/100 ml, Mg 1.5 mg/100 ml, and serum osmolality was 283 osmol/l. Electroencephalogram showed widespread arrhythmic delta activity over all regions and focal spikes over the left temporal area.

The patient's hospital course was complicated by pneumonitis and empyema requiring surgical drainage. He regained consciousness on the fifth day after admission to UCSF and was discharged 41 days later, without any neurological deficit.

DISCUSSION
An absence of oculocephalic or caloric-induced eye movements indicated structural or functional disruption of the vestibulo-ocular pathways (Klingon, 1952; McNealy and Plum, 1962; Plum and Posner, 1972). Patients who are comatose from drug intoxication (Bender et al., 1955; Nathanson et al., 1957) or metabolic encephalopathy (Silberpfennig, 1938; Plum and Posner, 1972) may lose vestibulo-ocular eye movements; recovery of these movements in those who survive is evidence against structural brain-stem damage. In patients who are comatose from other causes, absence of vestibulo-ocular eye movements has been cited as a sign of irreversible brain-stem damage and impending death (Rodríguez Barrios, 1966; Poulsen and Zilstorff, 1972).

Acute supratentorial lesions do not abolish vestibulo-ocular reflexes unless associated with transtentorial herniation and its brain-stem vascular complications (Vaernet, 1957; Needham et al., 1970). The absence of vestibulo-ocular reflexes in our patients occurred during stupor, not deep coma, and without other focal signs of brain-stem damage.

It is likely that subarachnoid haemorrhage contributed to stupor in case 1. According to Fisher (1969), loss of vestibulo-ocular reflexes cannot be attributed to subarachnoid haemorrhage alone. In experimental animals, the presence of blood in the ventricular fluid does not impair brain-stem function (Carpenter et al., 1967). Sambrook et al. (1973) reported that loss of vestibulo-ocular reflexes in patients with subarachnoid haemorrhage is invariably associated with deep coma, abnormal respirations, and absent pupillary responses. All of these patients died. Haemorrhage into the fourth ventricle in this circumstance is also fatal (McDonald, 1962; Ojemann and New, 1963). The patient in case 1 was not comatose, had normal vital signs, and recovered.

In the second case, loss of vestibulo-ocular reflexes was associated with superior sagittal sinus thrombosis. The possibility of coincident viral encephalitis could not be excluded. The aetiology of sinus thrombosis in this case may be related to the preceding viral illness (Krayerbühl, 1967) or dehydration subsequent to medical treatment.

Acute cerebral lesions may abolish the fast component of caloric nystagmus in the side opposite the lesion (Merwarth and Feiring, 1939). In monkeys this loss may persist for as long as one month (Pasik et al., 1961). Chronic cerebral
these combinations non-comatose present (Buchthal and Killam, 1966), stimulating the mesencephalic reticular formation of encephale isolé cats, observed either facilitation or inhibition of vestibular nystagmus. Collins (1963) stressed the varying effect of ‘alertness’ on the amplitude and duration of vestibular nystagmus in normal subjects. The fast phase of vestibular nystagmus is absent during sleep (Markham, 1972). A remote effect of bilateral hemispheric or midbrain dysfunction upon the vestibulo-ocular reflex arc, diachisis, may have contributed to the loss of these reflexes in our patients.

Barbiturates do not abolish vestibulo-ocular reflexes in conscious patients when used in therapeutic dosages. Serum phenobarbitone levels of 1–3 mg/100 ml are considered therapeutic, while levels greater than 3 mg/100 ml usually produce ataxia and nystagmus (Buchthal and Lennox-Buchthal, 1972a). Anderson et al. (1958) injected sodium amylobarbital intravenously into conscious normal subjects during caloric testing and observed transient loss of induced nystagmus. Barbiturate blood levels were not measured in their study. Rashbass and Russell (1961) observed no change in the fast or slow phases of vestibular nystagmus in similar experiments, although smooth pursuit movements became saccadic. The brain-stem reticular formation is extremely sensitive to barbiturates. Single cell microelectrode recordings in this area show depression of spontaneous and somatosensory-evoked activity even with small doses of pentobarbitone (Killam, 1962).

Therapeutic serum concentrations of diphenylhydantoin occur within a range of 1–2 mg/100 ml. Nystagmus and saccadic pursuit may be present with serum levels as low as 1 mg/100 ml (Buchthal and Lennox-Buchthal, 1972b). Absent vestibulo-ocular reflexes have been recorded in non-comatose patients intoxicated with combinations of diphenylhydantoin and phenobarbitone (Orth et al., 1967). Blood levels of these drugs in our patients were within the accepted therapeutic range during the period of vestibulo-ocular reflex loss.

Reversible loss of vestibulo-ocular reflexes in patients with structural cerebral hemispheric damage is extraordinary. Signs of transtentorial herniation were not present; therefore, we cannot attribute the loss of vestibulo-ocular reflexes in our patients to the cerebral lesions alone. We postulate that the combined effects of acute cerebral hemispheric damage and anticonvulsant drugs or subarachnoid blood acting upon the brain-stem substrate of the vestibulo-ocular reflex resulted in the loss of these reflexes.

On the basis of our cases, we conclude that the absence of the vestibulo-ocular reflexes in patients with acute cerebral lesions cannot be interpreted as pre-emptory evidence of irreversible brain-stem damage.

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