Suppression of brainstem reflexes in barbiturate coma

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Abstract
Brainstem reflexes were examined in 23 children treated with thiopentone infusion and correlated with serum thiopentone concentrations. The results suggest that if all brainstem reflexes are lost with a serum thiopentone concentration less than 40 mg/l, it is unlikely to be due to the thiopentone alone. Other causes including brain death need to be considered.

Results
The results are given in the table. Although other brainstem reflexes were frequently lost, the pupillary response to light was absent in only four patients. In two this coincided with the highest thiopentone concentrations recorded: in patient 1 it was 70 ml/g/l and in patient 2 it was 59 mg/l. The thiopentone concentration of patient 7 was 40 mg/l but a simultaneous clonazepam concentration was 1946 µmol/l (therapeutic range <15 µmol/l). In all three the light reaction returned on the next day. The fourth patient who lost all brainstem reflexes, patient 21, was brain dead at the time of the examination. The EEGs showed a good correlation with the thiopentone concentrations with electrocerebral silence present in patients 1, 7, and 8 (concentrations of 70, 40, and 38 mg/l), burst suppression in patient 10 (34 mg/l), and diffuse slowing with super added fast activity in patients 12, 22, and 23 (29, 9, and 5 mg/l). As this did not seem to be providing additional information formal EEGs were not done on the final 16 patients in the study.

Discussion
The dilemma over the diagnosis of brain death in patients treated with barbiturate coma would seem to be quite common as most sizable studies report a high mortality and three of our patients died. There is some information from these studies on the effect of barbiturates on brainstem reflexes. Lowenstein et al used pentobarbitone inducing a burst suppression pattern with concentrations ranging from 4 to 21 mg/l. All brainstem reflexes except the pupillary response to light were lost. Young et al also used pentobarbitone in four adults maintaining concentrations between 20 and 40 mg/l to produce a burst suppression pattern, again with preservation of the light reflex. Tasker et al noted that two of their patients lost the pupillary response to light at thiopentone concentrations of 58 and 77 mg/l respectively. The first patient recovered but the second died. It was observed with this patient that once the thiopentone had been stopped, a further five days of support were required before all the drug had been eliminated. Osorio and Reed used pentobarbitone to treat 12 adults and found that the pupillary light reflex was not observed when the serum pentobarbitone concentrations approached 50 mg/l. Dopamine infusion has been reported to cause fixed dilated pupils, but the dose was three times greater than the maximum dose used in our study and we feel this is unlikely to have been a contributing factor in our patients.

When faced with a patient who has lost all brainstem reflexes during barbiturate coma, one
The table is ordered from the highest to lowest thiopentone concentrations.

*A=absent; P=present; Pt=reaction to light present but sluggish.
**EThe outcome scale is 1=functionally normal; 2=mild handicap, functionally independent; 3=severe handicap requiring dependent care; and 4=died in intensive care unit.

approach is to wait for the drug to be fully metabolised before making a decision about brain death. This, however, may take many days and is an undesirable situation. The delay causes unnecessary suffering to the patient’s family and may adversely affect the morale of intensive care staff. Optimum conditions for organ transplantation may not be achieved and other children denied the benefit of the limited number of intensive care beds.

Our findings and those of the literature suggest that if all brainstem reflexes have been lost with a thiopentone concentration below 40 mg/l, the thiopentone alone is unlikely to be responsible. After other causes including intoxication with other drugs and local pathology producing fixed pupils have been excluded, the diagnosis of brain death should be considered. In the appropriate clinical setting, cerebral angiography can be performed without waiting for the drug to be fully metabolised.