The effect of thiopentone on somatosensory evoked responses and EEGs in comatose patients

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SUMMARY EEgs and somatosensory evoked responses from the brachial plexus, neck and scalp were recorded in seven comatose patients on continuous thiopentone infusion. Although pathological in five of the patients, the evoked responses were present in all. Additional amounts of thiopentone producing a full suppression of all spontaneous EEG activity had no effects either on the configuration of the evoked responses or on the central conduction times. This resistance of the somatosensory evoked responses to a deep and sustained thiopentone narcosis makes it a useful test in comatose patients receiving this treatment.

Evoked responses have proved valuable for determining the level and cause of coma and also to predict the final outcome of the comatose condition.1–12 Although the evidence of the value of massive barbiturate treatment in comatose patients has been conflicting,13 the treatment is in wide use and has complicated the interpretation of the evoked responses, since this drug has effects both on the visual14–17 and cortical somatosensory evoked potentials.14–18–25 Further, the clinical neurological examination of patients on continuous thiopentone infusion is of questionable value, since the drug effects can mimic most of the signs caused by coma of cerebral origin. The EEG is of minimal help since even small amounts of barbiturates have a profound effect on the records.26–28 It is therefore of value to determine whether any of the evoked responses are sufficiently resistant to thiopentone narcosis to be used as a clinical diagnostic test. In the present study the peripheral, cervical and cortical somatosensory evoked responses (SEP) were recorded and compared to the EEG records in seven comatose patients on thiopentone narcosis. No attempts were made to evaluate the prognostic value either of the SEP test or thiopentone narcosis. The purpose of the study was to test the sensitivity of the evoked responses and of the central conduction time to a relatively long lasting deep thiopentone narcosis so as to determine whether the SEP test could be reliably applied also in this group of intensive care patients.

Materials and methods

The seven patients examined in the present study were taken from a larger series of comatose patients treated in the neurosurgical intensive care unit.

Treatment with thiopentone was started mainly to prevent an expected rise in the intracranial pressure, and was initiated either when the patient was admitted to the intensive care unit or in a local hospital. The drug (thiopentone-Na 5 g/l in 5% glucose) was administered by continuous intravenous infusion. The infusion rate was guided by the cerebral perfusion pressure (CPP). Additional thiopentone was either given as several slowly administered bolus injections or by increasing the infusion rate. Serum levels of thiopentone were determined regularly several times a day. During the barbiturate treatment the patients were artificially ventilated and the PaCO₂ were kept within 3.5–5.4 kPa whenever possible. The intracranial epidural pressure (EDP) was constantly monitored with a minitransducer placed epidurally through a burr hole in the right parietal region.29

Arterial blood pressure was recorded with a fluid pressure transducer and together with the EDP continuously displayed on a multichannel pen recorder. An indication of the cerebral perfusion pressure (CPP) thus was available throughout the intensive care period. The mean arterial blood pressures were above 100 mm Hg and the cerebral perfusion pressures (CPP) were above 50 mm Hg in all patients when the SEPs and EEGs were recorded. The patients' temperatures were actively kept within 36–38°C throughout the intensive care period.

Recording and stimulation The EEG was recorded with...
an eight channel Elema Schöndering mingograph. Somatosensory evoked potentials (SEP) were simultaneously recorded from the brachial plexus (Erb's point), the neck at C5-level, and the scalp projection of the postrolandic hand area. The common reference electrode was placed on the forehead. The recording electrodes were commercially available Ag/AgCl skin electrodes. The analogue signals were recorded and simultaneously averaged on a Medelec MS 6 electrophysiological system.

The median nerves either in the wrist or in the elbow were used for stimulation, stimulation frequency was 3 Hz. Generally 300–500 analogue signals were averaged. High and low pass filters in the amplifiers were set to 16 Hz and 3-2 kHz respectively in all channels. The signal latencies were determined with an adjustable electronic marker and were always measured from the stimulus artifact to the peak negativity of the responses. The central conduction time was determined as the peak interval between the cervical (N13) and first negative cortical responses (N20). Central conduction times above 7-2 ms were regarded as abnormal, as were cortical response with an N20 amplitude less than 1 μV.

**Results**

EEG and somatosensory evoked responses (SEP) generally were taken on the second day after admission to the neurosurgical intensive care unit. The main data of the patients are given in the table. The EEG pattern as well as the corresponding thiopentone level varied considerably at this first examination (table). In three of the patients the EEG consisted of high voltage continuous slow wave activity while four patients had more or less pronounced "suppression-burst" pattern EEG consisting of periods with low voltage records interrupted by "bursts" of relatively high amplitude. The results failed to show any consistent relation between the thiopentone level and the individual EEG pattern (table) although increasing blood levels of the drug always reduced the EEG activity towards an isoelectric trace in individual patients. The somatosensory evoked responses recorded from the brachial plexus (N9), neck (N13) and scalp (N20) invariably were present in all patients. The cortical responses (N20) although present in all were of abnormally low amplitude (<1 μV) in four patients (table) and the central conduction time (N13–N20) prolonged (above 7-2 ms) also in four patients. The SEP test was normal in two patients (UH and AV) throughout their intensive care period. Two examples of the initial EEG records and SEP tests are given in fig 1 A–D. The corresponding thiopentone serum level are given above the respective EEG records (fig 1A, C).

Comparison of the SEP test abnormalities and the EEG records of the patients failed to show any consistent relationship (table). When additional thiopentone was administered, the effect on the EEG was rapid with slowing of the records and development of long suppression periods. The duration of the suppression periods varied from 3–7 minutes. When the drug administration was finished a "typical" suppression burst pattern with shorter periods of suppression (seconds) developed in all patients. The amount of thiopentone giving the long suppression periods varied considerably from patient to patient. SEP taken during the period with full suppression of all spontaneous EEG activity invariably failed to show any changes either in signal morphology or central conduction times (table). Two examples of SEP tests during full EEG suppression are given in fig 2 A–D. The records were

<table>
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<tr>
<th>Initials</th>
<th>Age (yr)</th>
<th>Diagnosis</th>
<th>First examination</th>
<th>After additional thiopental</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Serum thiopentone</td>
<td>EEG</td>
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<tr>
<td>TA</td>
<td>14</td>
<td>Head injury</td>
<td>87</td>
<td>Suppression</td>
</tr>
<tr>
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<td>12</td>
<td>Cardiac arrest</td>
<td>76</td>
<td>Delta/theta</td>
</tr>
<tr>
<td>EL</td>
<td>23</td>
<td>Head injury</td>
<td>212</td>
<td>Suppression</td>
</tr>
<tr>
<td>TF</td>
<td>17</td>
<td>Gun shot, brain injury</td>
<td>170</td>
<td>Delta/theta</td>
</tr>
<tr>
<td>OT</td>
<td>34</td>
<td>Head injury</td>
<td>118</td>
<td>Suppression</td>
</tr>
<tr>
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<td>9</td>
<td>Strangulation</td>
<td>136</td>
<td>Theta/delta</td>
</tr>
<tr>
<td>UH</td>
<td>21</td>
<td>Massive intraventricular haemorrhage from AVM</td>
<td>124</td>
<td>Suppression bursts</td>
</tr>
</tbody>
</table>

*Pathologically prolonged values (above 7-2 ms).
†Indicates which patients had pathological cortical responses (N20 lower than 1 μV, or abnormal response configuration).
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The EEGs (A, C) and SEP records (B, D) from two of the patients taken after 48 hours of deep thiopentone narcosis. The actual thiopentone serum levels are given above EEG records in A and C. Time scale for the EEG records (1 s between vertical lines) is given in the uppermost records in A and C. The records in B and D represent plexus (lower), cervical (middle) and cortical (upper) evoked responses, Time scale in ms.

Discussion

Previous human and experimental studies have demonstrated that barbiturate anaesthesia abolishes the later components of the cortical somatosensory evoked response while having relatively little effect on the earlier waves. The data were confirmed in the present study, showing that the first negative wave (N20) of the cortical response as well as the cervical (N13) and peripheral plexus (N9) evoked responses were present in all patients on sustained thiopentone narcosis. The N20 of the cortical response probably represents the first postsynaptic response to the thalamo-cortical volley, while the cervical evoked response evidently consists of synaptic as well as non-synaptic subcomponents generated in the cervical medulla. N9 represents the peripheral action potentials in the brachial plexus fibres. The fact that deep thiopentone narcosis had no effect on either of these responses or the "conduction time" between them, thus indicates that impulse transmission in the fast conducting dorsal column-lemniscal afferent pathway was resistant to large amounts of thiopentone. Corresponding results from human and animal data have been previously reported. The mechanism for the relative "insensitivity" of this system to barbiturates is not clear, but it may be related to the organisation of this particular afferent system with the few and powerful excitatory synapses involved. The effect of barbiturates on the central nervous system is mainly inhibitory, the inhibition partly being produced by enhancing the effect of inhibitory synapses. Barbiturates also have a more pronounced
inhibitory effect on multisynaptic pathways than on pathways with one or few synapses involved.\textsuperscript{21} 34-40 Since no EEG records were taken prior to thiopentone administration, it was not possible to decide to what extent the prevailing "brain injury" contributed to the initial EEG patterns. The fact that the SEP test was pathological in five of the patients examined, either with a reduced amplitude of N20 or with a prolonged central conduction time (N13-N20), however, evidently reflected the presence of a brain injury or ischaemia rather than thiopentone narcosis. The two patients (AV and UH, table) with normal responses and conduction times throughout their intensive care periods as well as the general lack of effect on the SEP test in all patients when additional thiopentone was administered further gave support to this assumption. The present data also demonstrated that amounts of thiopentone sufficient to suppress fully all spontaneous EEG activity for several minutes had no effect on the somatosensory evoked responses or conduction times. Whether the responses would remain unchanged during a longer period with isoelectric EEG is unknown. However, Trojaborg and Jørgensen\textsuperscript{11} have shown that the cortical somatosensory evoked response may persist and be well preserved (and indicate a relative good prognosis) in comatose patients with a totally isoelectric EEG. Furthermore, experimental data indicate that no qualitative physiological or biochemical changes occur in the brain at anaesthetic levels sufficient to produce a sustained isoelectric EEG.\textsuperscript{41-43} Massive doses of thiopentone producing long lasting isoelectric EEG have failed to reveal any toxic effects on the central

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Fig 2  The EEGs (A, C) and SEP records (B, D) of the same two patients as in fig 1 after additional amounts of thiopentone. The actual thiopentone serum levels are given above EEG records in A and C. Time scale for the EEG records (1 s between vertical lines) is given in the uppermost records in A and C.
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nervous system and have been used with some success in intensive care medicine. It is therefore unlikely that even a long lasting thiopentone induced suppression of spontaneous EEG activity would produce any major changes in the somatosensory evoked responses. Evidence favouring this hypothesis is further given by experimental data showing that although spontaneous unit activity may be fully depressed during barbiturate anaesthesia, the neurons were still able to respond to afferent stimuli. 

Clinical and prognostic evaluation of patients in thiopentone narcosis is generally difficult, being limited by all the restraints encountered when patients have to be examined while in deep narcosis. EEG offers little help in this situation other than being a monitor of the anaesthetic level. Moreover, the variability in this relation between the individual EEG changes and the actual serum level of thiopentone makes the judgement even more complex.

Brain stem evoked potentials (BAER) have been used with some success in comatose patients and are relatively resistant to anaesthetics. However, Goldie et al have reported that 52% of the patients in coma had normal BAER, and in general there was no correlation between the BAER of these patients and the clinical outcome. Visual evoked responses are unreliable in patients on thiopentone narcosis since the response morphology as well as the latencies are subject to changes during anaesthesia.

In conclusion we therefore feel that the present somatosensory evoked responses, including the central conduction times, offer a valuable test in evaluating the comatose intensive care patients in thiopentone narcosis.

References


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