Prediction of outcome in severe head injury based on recognition of sleep related activity in the polygraphic electroencephalogram

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Abstract
This study shows that the continuing presence of activity similar to normal sleep in the EEG in conjunction with the EEG polygraph (EEGP) can be used to determine the severity of brain damage after head injury. Recordings were taken within seven days of head injury from 154 unselected patients after resuscitation and emergency surgery. Sixteen patients with ongoing seizures were excluded. In the remaining 138 patients the presence of activity in the EEG, EEGP, or both, which can also be recognised in normal alertness and sleep, was noted. Particular attention was paid to the presence or absence of arousal related phasic activity involving EEG, motor, and autonomic changes. The traces were allocated to one of five groups: group 1, wakeful traces with normal alpha in at least one hemisphere; group 2, sleep-like traces with K complexes responsive to stimulation; group 3, traces with phasic activity related to abnormal spontaneous arousal including EEG changes; group 4, traces with abnormal spontaneous arousal activity without EEG changes; group 5, traces with no spontaneous arousal activity. The mean follow up was 21.5 months.

Comparison between the EEG/EEGP findings and the Glasgow coma scale at the time of the recording showed the EEG/EEGP to be the better predictor of outcome, particularly for individual patients.

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Keywords: head injury; outcome; sleep/wake mechanisms; EEG polygraph; Glasgow coma scale

At the onset of sleep there are well established changes from the wakeful alpha dominant EEG to the patterns of stages 1 to 4 sleep. More recently, spontaneous changes in the level of arousal have been shown both in normal sleep and in brain damage, which are related to the microstructure of sleep. These studies report spontaneous rhythmic changes in the arousal level of the EEG at intervals between seven and 60 seconds that are associated with autonomic and motor changes. It has been suggested that these phasic changes are related to a physiological arousal cycle at sleep onset and that the same mechanism is responsible for the phasic activity seen in coma. There is evidence to support this concept from animal experiments concerning the arousal and sleep/wake mechanisms.

This paper shows that physiological EEG/EEG polygraph (EEGP) activity related to sleep is recognisable in the damaged brain, correlates well with the severity of the damage after head injury, and can be used to predict outcome.

Patients and methods

Electroencephalograms, EEGPs, or both were obtained from 154 unselected patients (113 male, 41 female), admitted with head injury to the neurosurgical unit at The Brook General Hospital. Their ages ranged from 3 to 72 years (mean age 27.1 indicating a high proportion of younger patients). The clinical management included early resuscitation, often with an elective period of assisted ventilation using a muscle relaxant where necessary. High dose barbiturate treatment was not used. All electrophysiological traces were obtained within seven days of the injury and after resuscitation and emergency surgical procedures, using an SLE E8b or 100T machine at the bedside with silver/silver chloride surface or platinum intradural needle electrodes situated according to the modified Maudsley system of electrode placement. For the examples given in the figures, unless stated otherwise, silver/silver chloride disc electrodes were used, gain 50 μV/cm; low frequency filters 0.5 Hz; high frequency filters 70 Hz. In 18 patients the EEG alone was recorded. In 63 patients polygraphic records were obtained which included an ECG recorded from the shoulders, a surface EMG from a suitable muscle, and respiration from a thermistor in the nostril or the expiratory valve of the ventilator. The ECGs were
analysed by measuring the R wave interval to determine the beat to beat heart rate and plotting the results at one second intervals to produce a heart rate graph. In 73 patients the autonomic variables were obtained from a Hewlett Packard cardiorespiratory monitor (modification of a foetal heart rate monitor).

The recording methods were designed to provide an assessment of the physiological state of the patient coincident with the clinical assessment and were therefore not prolonged, lasting between 30 and 60 minutes. The traces were always observed during the recording by the same electroencephalographer (BME). Particular care was taken to ensure that the patient achieved the highest level of general arousal possible by stimulation of varying degrees of severity from calling the name to deep pain; the effects of this on the EEG polygraph were noted. Changes in the level of arousal, either spontaneous or after stimulation, were identified from the EEG and polygraph. Activity was considered to be related to arousal if it could be produced, altered, or when spontaneously present, reproduced, by stimulation given at the lower arousal levels. Clinical details, including the Glasgow coma score on a 14 point scale, were noted by the clinical team at the time of the record.

SEIZURE GROUP
Sixteen patients had continuous epileptiform activity, which made arousal changes impossible to assess by EEG. These patients were excluded from further study.

The remaining 138 patients, without seizures, formed the study group. These were divided into five subgroups determined by the EEG and polygraphic findings. The principal criteria were the level of general arousal obtainable and the changes in arousal produced in any part of the EEG/EEGP, either spontaneously or after stimulation. When the EEG was asymmetric the findings from the more normal hemisphere were used for grouping.

Group 1: wakeful records (fig 1)
This group could be recognised from the EEG alone. There was a well formed and sustained postcentral alpha rhythm for at least part of the time over one or both hemispheres. The distribution and reactivity of the alpha and the presence of eye movements distinguished the findings from alpha coma.

The eye movements were characteristic, either blinking or rolling.

The polygraph did not contribute further useful information.

Group 2: sleep-like records (fig 2)
This group could be recognised from the EEG alone. The traces showed stage 2 sleep activity with K complexes, which could be reproduced by stimulation, on a background of low voltage theta (4–7 Hz) and delta (< 4 Hz). Records with only spindle activity or
Figure 2. Group 2: sleep-like trace. A boy aged 11 was unconscious from the time of injury. At the time of admission he had flexor arm movements, groaned, but did not open his eyes (Glasgow coma scale grade 6). His right pupil was fixed and constricted. He deteriorated after admission, with fixed dilated pupils, bilateral extensor movements, and no vocalisation (Glasgow coma scale grade 4); CT showed a film of extradural blood, without mass effect, for which surgery was not considered necessary. He was ventilated electrically and made a good recovery within 48 hours with a transient left hemiparesis. The EEG on the day after injury (platinum needle electrodes) shows a background of low voltage theta and delta with K complexes following minimal stimulation (S), firstly by blowing on the skin and then by calling the name.

potentials resembling K complexes that were not sensitive to stimulation were not included.

Eye movements were present, unless pharmacologically paralysed, but were not contributory.

The polygraph added no useful information.

Group 3: abnormal spontaneous arousal activity with EEG change (fig 3A)
The characteristic findings were changes in the arousal level, both spontaneous and after stimulation. These involved all parts of the polygraph trace including the EEG.

Eye movements were present, unless pharmacologically paralysed, but were not contributory.

Polygraphic information was essential to distinguish this group from groups 4 and 5. Changes in arousal were accompanied by increases in beat to beat heart rate, respiration rate, and muscle activity, which could be phasic or irregular, and were often exaggerated (fig 3A), with changes in heart beat of up to 100 beats/min within a few seconds.

Group 4: abnormal spontaneous arousal activity without EEG change (fig 3B)
The criteria for this group were the same as those for group 3 except that arousal changes were only present in the other parts of the polygraph and were not seen on the EEG.

Group 5: no spontaneous arousal activity (fig 4)
This group showed no spontaneous arousal activity during the period of the recording (30–60 minutes). Changes on the EEG were never seen.

Spontaneous eye movements were absent even when the patient was not paralysed.

Changes in heart beat were not seen spontaneously (apart from respiratory sinus arrhythmia) but stimulation sometimes produced a brief poorly sustained response in heart beat, EMG, or respiration (fig 4A).

Outcomes were obtained from the clinical notes; all patients were assessed at three months. Among the survivors, those in unremitted coma were classified as vegetative, those completely recovered were discharged, and those with any disability were followed up for varying times between six and 84 months. Outcomes were graded according to the criteria of Jennett and Bond as (1) good recovery, (2) mild disability, (3) severe disability, (4) persistent vegetative state, (5) death. These groups were reduced to three: good 1 and 2, disabled 3, poor 4 and 5.

Statistical analysis of the relation between the clinical outcome and the EEG/EEGP groups was achieved by a $\chi^2$ (8 df) test.

Results

OUTCOMES
Table 1 shows the outcomes for the study group. Sixty six patients died within three months of their injuries. Follow up in the 72 survivors was between three and 84 (mean 21.5) months. There were 55 (40%) good outcomes of whom 33 were back to normal and 23 were mildly disabled. Thirteen (9%) were severely disabled. Seventy (51%) poor outcomes comprised 66 deaths and four vegetative patients.
Figure 3  Group 3: abnormal spontaneous arousal activity. (A) With EEG change. A girl aged 16 was unconscious from the time of a road accident. Her pupils were equal and reactive. She responded to pain by flexion of the right and extension of the left arm. She made no sound but occasionally opened her eyes (Glasgow coma scale grade 7); CT showed some generalised swelling. She was electively ventilated for 48 hours. Her recovery was slow but after five months there was no residual deficit. EEG was on the day after injury. Above: EEGP, channels 1–6 EEG, channel 8 ECG. Below: cardiorespiratory monitor, respiration from ventilator expiratory valve (functioning poorly). The EEG shows bursts of high voltage slow waves both spontaneously and on painful stimulation. The monitor shows very pronounced and irregular beat to beat heart rate changes spontaneously and on stimulation (X on both traces). (B) Without EEG change. A man aged 41 was admitted after falling off his motor cycle. On admission he was localising pain with his right arm, his eyes opened to pain, and he groaned occasionally (Glasgow coma scale grade 8). His right pupil was fixed and dilated and the left pupil reacted; CT showed bispinal and occipital fractures with left and right parietal and frontal and right temporal contusions; also a small right frontotemporal intracerebral clot. He eventually made a good recovery with some memory impairment (last seen six years later). The traces were taken two days after injury. Above: EEGP, channels 1–8 EEG, channel 9 ECG. Below: cardiorespiratory monitor. The EEG showed no change when the patient was aroused spontaneously. The monitor shows periodic respiration with accompanying beat to beat heart rate changes occurring at about 30 second intervals; X marks the spontaneous arousal movement on both traces.
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Figure 4  Group 4: no spontaneous arousal activity. (A) With EEG activity present. A man aged 44 fell off a ladder on to his head. On admission there was no movement in his legs but his arms extended; he was silent with closed eyes (Glasgow coma score grade 4). Both pupils were fixed and dilated. He was treated with dexamethasone and intravenous fluids; his arms then responded purposefully (Glasgow coma score grade 6) and his pupils became sluggishly reactive. A skull radiograph showed an occipital fracture. Over the next four days he deteriorated and died. Postmortem showed extensive contusion and petechiae throughout the hemispheres, cerebellum, and brain stem. Traces were obtained on the second day after injury. Above: EEGP, channels 1–6 EEG, channel 7 respiration, channel 8 ECG. Below: cardiorespiratory monitor. The EEG shows a mixture of frequencies from 2–7 Hz with no response to painful stimulus (sternal pressure). The monitor shows almost unvarying respiratory sinus arrhythmia with minimal change to some stimuli; X marks the moment of stimulation on both traces. (B) Without EEG activity. A girl aged 12 was knocked down by a car. On admission she flexed both arms. She was silent with closed eyes (Glasgow coma score grade 5). Her pupils reacted sluggishly to light; CT showed severe fractures of the left parietal and frontal bones with brain swelling, worse on the left and shift of the midline structures. She was electively ventilated and her wound explored. The skull fragments were mobile and there was necrotic extruded brain. She did not improve and died two days later. Traces were obtained on the day of injury, after surgery while the patient was artificially ventilated. Above: EEGP, channels 1–5 EEG. Below: cardiorespiratory monitor. The EEG shows only some muscle and ECG artefact. The EEG and beat to beat heart rate did not respond to a painful stimulus (X). The heart beat trace shows only slight undulations, not related to respiration.
Table 1 Outcome in the study of 138 seizure free patients compared with a multicentre study in 1976

<table>
<thead>
<tr>
<th>Group</th>
<th>Died</th>
<th>Persistent vegetative state</th>
<th>Severe disability</th>
<th>Moderate disability</th>
<th>Good</th>
</tr>
</thead>
<tbody>
<tr>
<td>EEGP 1976</td>
<td>66 (48%)</td>
<td>4 (3%)</td>
<td>13 (9%)</td>
<td>23 (17%)</td>
<td>32 (23%)</td>
</tr>
<tr>
<td>ASA</td>
<td>49%</td>
<td>3%</td>
<td>10%</td>
<td>17%</td>
<td>20%</td>
</tr>
</tbody>
</table>

ASA = abnormal spontaneous arousal; NSA = no spontaneous arousal.

Table 2 Glasgow coma scale in relation to the EEG/EEGP groups

<table>
<thead>
<tr>
<th>Scale</th>
<th>Wakeful</th>
<th>Sleep-like</th>
<th>ASA/EEG</th>
<th>NSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; 8</td>
<td>8 (89%)</td>
<td>4 (27%)</td>
<td>6 (10%)</td>
<td>0</td>
</tr>
<tr>
<td>6,7,8</td>
<td>1 (11%)</td>
<td>9 (60%)</td>
<td>34 (59%)</td>
<td>7 (12.5%)</td>
</tr>
<tr>
<td>&lt; 6</td>
<td>0</td>
<td>2 (13%)</td>
<td>18 (31%)</td>
<td>49 (87.5%)</td>
</tr>
</tbody>
</table>

ASA = abnormal spontaneous arousal; NSA = no spontaneous arousal.

Table 3 Relation between the Glasgow coma scale and final outcome

<table>
<thead>
<tr>
<th>Outcome</th>
<th>&gt; 8</th>
<th>6, 7, 8</th>
<th>&lt; 6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Good</td>
<td>10 (55%)</td>
<td>34 (67%)</td>
<td>11 (16%)</td>
</tr>
<tr>
<td>Disabled</td>
<td>2 (11%)</td>
<td>5 (10%)</td>
<td>6 (9%)</td>
</tr>
<tr>
<td>Poor</td>
<td>6 (33%)</td>
<td>12 (23%)</td>
<td>52 (75%)</td>
</tr>
</tbody>
</table>

GLASGOW COMA SCALE SCORES

Table 2 shows the Glasgow coma scale scores for the study group: > 8 18 (13%) patients. 6, 7, 8 51 (31%) patients, < 6 69 (50%) patients. Table 3 shows the relation of the Glasgow coma scale grades to outcome.

EEG/EEGP GROUPS

Group 1: Wakeful records
There were nine (6%) patients (two EEG, seven EEGP), average age 50 in this group (fig 1).

Six EEGs were asymmetric, five of which were among the patients with poor outcomes. Four of these patients had intracranial haematomas or haemorrhagic contusions, as did the one disabled patient.

The outcomes were: good three (33%), disabled one (11%), and poor five (55%). The findings were of no predictive value (table 4).

All were in Glasgow coma scale grades 8 or higher (table 2).

Group 2: Sleep-like records
The group contained 15 (11%) patients (four EEG 11 EEGP), average age 15 (fig 2).

The outcomes were: good 14 (93%), disabled one (7%), poor none. This EEG pattern was strongly associated with a good recovery, (table 4).

There was a wide spread of Glasgow coma scale grades with two patients in grade 4 and three in grade 6 (table 2).

Group 3: Abnormal spontaneous arousal activity with EEG change
There were 44 (32%) patients (all EEGP), average age 23 (fig 3A).

The EEG activity varied from patient to patient. Higher arousal levels were accompanied by bursts of high voltage delta activity whereas the lower arousal periods contained either a mixture of theta and delta or delta of much lower amplitude. High levels of arousal were common and it was often necessary to wait for many minutes before seeing a period of low arousal. In many cases there was asymmetry of the delta which usually correlated with asymmetries in the clinical state.

The arousal episodes often occurred rhythmically with a periodicity of between 20 and 60 seconds but they could also be irregular (fig 3A). They could be brief or prolonged, sometimes lasting up to 10 to 15 minutes.

Outcomes in group 3 were: good 33 (75%) patients, disabled four (9%) patients, poor seven (10%) patients. These findings were associated with a good recovery but less so than in the sleep-like group (table 4).

Group 4: Abnormal spontaneous arousal activity without EEG change
This group contained 14 (10%) patients, average age 24 years. All had EEGP (fig 3B). The EEG showed continuous delta activity, sometimes of low voltage, and often mixed with theta and alpha. Some patients showed spindles of beta (two of whom made good recoveries). The traces often showed asymmetry of the delta activity.

Arousal activity similar to that seen in group 3 was present in the polygraph traces other than the EEG, either rhythmic (fig 3B) or irregular.

The outcomes for this group were: good four (29%), disabled two (14%), poor eight (57%). This finding was of no clear predictive value.

Groups 3 and 4 included most Glasgow coma scale grades: > 8 six patients (10%), 8-6 34 patients (59%), < 6 18 patients (31%) (table 2).

Group 5: No spontaneous arousal activity
This group contained 56 (41%) patients (five EEG, 54 EEGP), average age 27 (fig 4).

The EEG findings varied. Isoelectric records were seen in 23 patients. Continuous unvarying delta, often of low voltage, sometimes mixed with theta and alpha frequencies, and occasionally spindles of beta (not including the patient who recovered) were seen in 24 patients. Nine patients showed alpha coma.

The outcomes were good one (2%), disabled five (9%), poor 50 (89%). This finding was strongly associated with a poor outcome (table 4).

Only the lower Glasgow coma scale grade were represented in this group: > 8 0, 8-6 seven (12.5%), < 6 49 (87%) (table 2).

Table 4 Relation between the five EEG/EEGP groups and final outcome (square brackets show predicted outcomes)

<table>
<thead>
<tr>
<th>Group</th>
<th>Wakeful</th>
<th>Sleep-like</th>
<th>ASA + EEG</th>
<th>ASA – EEG</th>
<th>NSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Good</td>
<td>3 (33%)</td>
<td>14 (93%)</td>
<td>33 (75%)</td>
<td>4 (29%)</td>
<td>1 (2%)</td>
</tr>
<tr>
<td>[5-9]</td>
<td>[5-98]</td>
<td>[17-54]</td>
<td>[5-58]</td>
<td>[5-32]</td>
<td></td>
</tr>
<tr>
<td>Disabled</td>
<td>1 (11%)</td>
<td>1 (7%)</td>
<td>4 (9%)</td>
<td>2 (14%)</td>
<td>5 (9%)</td>
</tr>
<tr>
<td>[0-9]</td>
<td>[1-41]</td>
<td>[1-14]</td>
<td>[1-32]</td>
<td>[5-27]</td>
<td></td>
</tr>
<tr>
<td>Poor</td>
<td>5 (55-5%)</td>
<td>0</td>
<td>7 (16%)</td>
<td>8 (57%)</td>
<td>50 (89%)</td>
</tr>
<tr>
<td>[4-56]</td>
<td>[7-61]</td>
<td>[22-31]</td>
<td>[7-10]</td>
<td>[28-4]</td>
<td></td>
</tr>
</tbody>
</table>

\[ r^2 = 0.52 (8 df), P < 0.001. \]

Good = mild disability + normal; Poor = died + vegetative; ASA = abnormal spontaneous arousal; NSA = no spontaneous arousal.
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STATISTICAL RESULTS
Table 4 shows the results analysed by \( \chi^2 \) test (8 df). The results were highly significant.

Discussion
This study shows that the physiological EEG/EEGP changes of normal sleep and wakefulness can be detected in head injury and can be used to provide information about the severity of the brain damage and hence predict the eventual outcome.

There is increasing evidence to suggest that the EEG/EEGP activity of sleep onset is concerned with the active change in behavioural state from wakefulness to delta sleep. The findings reported here suggest that sleep and waking may result from the operation of an intracerebral system, analogous to the pyramidal or visual systems, damage to which results in malfunction. The physiological mechanisms involved can still be identified in the damaged brain.

Of particular importance in this context is the experimental work on sleep spindles. In normal sleep, alternations between high and low arousal at intervals of seven to 60 seconds, associated with autonomic and motor changes are recognised. These are thought to relate to the microstructure of sleep as opposed to the macrostructure of the circadian and ultradian rhythms. Similar activity has long been recognised in the damaged brain with varying periodicities between 15 and 60 seconds. Many studies have linked this arousal related activity to changes in autonomic or cerebral autoregulatory mechanisms such as cerebral blood flow or CSF pressure. Although the phasic alternations in the damaged brain are not identical with those in normal sleep, particularly in respect to the type of EEG activity and the magnitude of the autonomic changes, there is little doubt that they are the same phenomenon. It is these phasic changes that relate to the abnormal spontaneous arousal activity described in groups 3 and 4.

The five EEG/EEGP trace types described show increasing levels of abnormality from wakefulness to absence of all arousal activity. The traces have been grouped in relation to the general level of arousal achievable, and the responsiveness of the EEG/EEGP to stimulation, rather than any particular EEG appearances. For this reason spindle activity alone, (Courjon’s spindle coma) was not used to categorise the sleep-like group (2) in the absence of reactive K complexes.

The EEG/EEGP findings have been used to determine the neurophysiological state of patients at the time of the examination in a similar way to the clinical assessment provided by the Glasgow coma scale. The Glasgow coma scale is an effective method of assessing patients after head injury. These clinical tests record the type of motor response and level of awareness (the ability to relate appropriately to the surroundings) at maximum arousal. These are similar physiological functions to those shown by the EEG/EEGP, although the EEG/EEGP includes additional information about the quality of cortical function and the presence or absence of spontaneous arousal activity. To emulate the universal application of the Glasgow coma scale there was no attempt to select cases on the basis of injury type, age, or recent surgery.

The outcomes from the study group of 138 patients show the group to be of similar composition to that in the multicentre study undertaken in 1976 (table 1). The Glasgow coma scale is known to provide one of the best correlations with outcome in head injury. In some circumstances, however, it can be difficult to perform, particularly in patients being ventilated, with severe facial injuries, or with peripheral palsies or fractures. Therefore an additional method of assessment is of practical clinical importance. It is suggested that the EEG/EEGP provides such a method.

Table 3 shows the outcome of the study group in relation to the Glasgow coma scale. There is a correlation between the coma grades < 6 and a poor outcome, the middle grades 8–6 show a mixed result with a preponderance of good outcomes. The highest coma grades did not correlate as well with a good outcome as other studies. Patients with high Glasgow coma scale grades were not usually admitted to the neurosurgical unit so that this finding probably reflects case selection by the admitting physicians. Some had a hemiparesis and were admitted to exclude a treatable cause for this, others had traumatic subarachnoid haemorrhage requiring investigation.

EEG/EEGP GROUPS (TABLE 1)
Group 1: the wakeful group
Despite a high level of awareness this group did not show a relation with good outcome. The asymmetries in the EEGs of six patients suggested that they had one severely damaged hemisphere and five had haematoma or haemorrhagic contusions. This finding probably reflects the case selection referred to and accounts for the poor outcome. The severity of damage to one hemisphere dictates the outcome here. As these patients were in Glasgow coma score grades 8 or above they represent some of the poor results among the higher Glasgow coma score grades shown in table 3.

Group 2: the sleep-like group
These patients showed responsive EEG activity associated with normal stage 2 sleep. This group showed a strong positive correlation with good outcome. Patients did not always have high Glasgow coma score scores; two were in grade 4 and three in grade 6. The excellent prospects for this group account for some of the patients with good outcomes among the lower Glasgow coma score grades (table 4). The relation of these EEG findings to good outcome has been shown several times before.
Groups 3 and 4

Both of these groups showed abnormal spontaneous arousal activity. The EEG/EEGp showed spontaneous alternations in arousal level, which could be duplicated by stimulation. There were associated, often exaggerated, autonomic and motor changes. The difference between groups 3 and 4 related solely to the presence or absence of EEG changes after arousal. These two groups could not be identified from the EEG alone. Some polygraphic information was essential, the most valuable being the beat to beat heart rate. The addition of the cardiorespiratory monitor was of great value in facilitating the distinction between the two abnormal spontaneous arousal groups (3 and 4) and group 5 (no spontaneous arousal). Groups 3 and 4 together had a similar outcome profile as the patients with Glasgow coma scale scores 6–8 (tables 2 and 3).

For group 3, the abnormal spontaneous arousal with EEG change is probably the same as the cycling EEG pattern that has been recognised for many years.3,5,20,24 Group 3 was associated with a good outcome (75% good) but less so than the sleep-like group.

For group 4 the arousal activity could only be recognised by autonomic or motor activity. Although the results were better than those of group 5 only four (29%) patients made good recoveries. Group 4 contained some traces that may have been allied to the sleep-like activity of group 2; traces with beta spindle but no reactive K complexes were excluded from group 2 and relegated to group 4. This is the probable explanation for the two rapid and complete recoveries mentioned in the results section.

For group 5 there was no evidence of spontaneous arousal. Only one patient made a good recovery and the group is strongly associated with a poor outcome. The group included a high proportion of the patients with low Glasgow coma scale scores (table 3). Table 4 shows 11 (16%) unexpectedly good recoveries in patients with Glasgow coma scale scores < 6, a finding also recorded in other studies.9,10 The EEG/EEGp can identify surviving normal cerebral, autonomic, and motor activity,18 and thus provides a better prediction of outcome than the Glasgow coma scale alone.

The EEG findings in this group were varied. There were isoelectric records, records with continuous unvarying delta activity, and nine cases of alpha coma. Some mixture of frequencies was often present and several traces had spindles of fast activity, showing that this finding is not in itself a predictor of good outcome.

If the wakeful group (1) is excluded, the statistical studies show a steady deterioration of outcome across the other four groups, which reflects the increasing damage to cerebral function (table 4).

The relation of poor outcome to increasing age in head injury is well established10,12 and the EEG/EEGp groups described have differing age profiles. Whereas groups 3 and 4 (abnormal spontaneous arousal with and without EEG changes) and group 5 (no spontaneous arousal) were similar to the mean ages of the study group as a whole group 1 (wakeful) had a much higher average age (50 years). It is possible that older brains respond to injury differently from younger ones, possibly as a result of primary or secondary vascular changes such as infarction of haemorrhage. This is supported by the finding of six unilateral hemispheric lesions among these patients. Group 2 (sleep-like) had a much lower average age (15 years) than the study group as a whole. The EEG appearances here were those of normal sleep implying no serious degree of cerebral harm. A possible explanation for this finding may be a greater susceptibility in younger patients to reversible brain stem shock, which has been postulated by some workers.

The findings show the value of the EEG/EEGp in the evaluation of patients with head injury, both in its own right and to provide supplementary information when the Glasgow coma score is difficult to use. The EEG/EEGp gives a better picture than the Glasgow coma scale of the functional state of the neuraxis in individual patients.

We thank Professor B Everitt for statistical advice.
Coca shrub (*Erythroxylum coca*)

The main alkaloid found in the coca leaves is cocaine. For centuries the Indians of Peru and Bolivia chewed the leaves for their stimulating effects. In the middle of the 19th century the linguist J J Von Tschudi became interested in the leaves for increasing physical performance. In Paris a manufacturer named Mariani made wine from coca leaves and marketed it as a tonic. This became very popular in Europe and the United States. Enthusiasts included President William McKinley, Thomas Edison, and the Tsar of Russia and Mariani received a medal of appreciation from the Pope. At various times attempts were made to use coca as a treatment for syphilis and whooping cough but without obvious success. In 1859 the Austrian explorer and ship's doctor Karl Von Scherzer brought dried coca leaves to Europe and gave them to the German chemist Wöhler of Göttingen for analysis. In 1859 Wöhler's pupil Niemann succeeded in extracting the effective component of the leaves, which he called cocaine. Moreno y Maiz, a Peruvian doctor, wrote, in 1868, of cocaine's ability to abolish sensation. He posed the question as to whether cocaine could be used as a local anaesthetic.

In 1884, after using it to treat his own depression, Sigmund Freud introduced cocaine to the physicians of Vienna. He maintained that it was valuable for that disorder, for eliminating nervous stomach complaints, and for augmenting mental and physical efficiency. He also commented on its ability to render mucous membranes insensitive. Freud attempted to cure a variety of nervous diseases with cocaine, even hydrophobia, but failed. In 1885 his treatment of a patient with trigeminal neuralgia by injection of cocaine was unsuccessful. In all probability he missed the nerve.

In 1884, Dr. Carl Koller, a colleague of Freud, discovered that the human eye could be rendered insensitive to pain with cocaine, so heralding the start of local anaesthesia. Other men took up and advanced the idea of local anaesthesia, bringing cocaine out of the restricted field of ophthalmology. Halsted injected cocaine into the inferior alveolar nerve; a discovery that revolutionised dentistry. Halsted later became addicted to the drug; he treated his addiction with morphine and became a morphine addict. Earlier, in a reverse approach, Freud had treated his friend Ernst Fleish's morphine addiction with cocaine and converted him into a cocaine addict. Sir Arthur Conan Doyle had his fictional detective, Sherlock Holmes, take cocaine to keep his wits occupied when not working on a case.

In 1886 John Pemberton of Atlanta, Georgia, introduced Coca Cola, originally an elixir from the cola leaves and caffeine rich extracts from the cola nut. He promoted it as a headache remedy and stimulant. Cocaine was removed from the formula in 1903.

Local anaesthetics such as lignocaine, benzocaine, and procaine have cocaine's anaesthetic properties without its stimulatory side effects.

The coca shrub was shown on a stamp issued by Rwanda in 1969 (Stanley Gibbons 311, Scott 301).