The combined monitoring of brain stem auditory evoked potentials and intracranial pressure in coma. A study of 57 patients

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Abstract
Continuous monitoring of brainstem auditory evoked potentials (BAEPs) was carried out in 57 comatose patients for periods ranging from 5 hours to 13 days. In 53 cases intracranial pressure (ICP) was also simultaneously monitored. The study of relative changes of evoked potentials over time proved more relevant to prognosis than the mere consideration of “statistical normality” of waveforms; thus progressive degradation of the BAEPs was associated with a bad outcome even if the responses remained within normal limits. Contrary to previous reports, a normal BAEP obtained during the second week of coma did not necessarily indicate a good vital outcome; it could, however, do so in cases with a low probability of secondary insults. The simultaneous study of BAEPs and ICP showed that apparently significant (> 40 mm Hg) acute rises in ICP were not always followed by BAEP changes. The stability of BAEP’s despite “significant” ICP rises was associated in our patients with a high probability of survival, while prolongation of central latency of BAEPs in response to ICP modifications was almost invariably followed by brain death. Continuous monitoring of brainstem responses provided a useful physiological counterpart to physical parameters such as ICP. Serial recording of cortical EPs should be added to BAEP monitoring to permit the early detection of rostrocaudal deterioration.

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Patients and methods
a) BAEP monitoring system
The monitoring system used for this study has been described in detail elsewhere. Briefly, it consists of a one-channel standard EP device coupled to a microcomputer (IBM-PC) which controls all operations. The recording electrodes are either Ag/AgCl disks fixed with collodion jelly or sterilised subcutaneous needles placed at Cz (positive input of the preamplifier) and over the mastoid process ipsilateral to the stimulated ear (negative input). A third electrode over the contralateral mastoid process serves as ground. The stimuli, delivered at a rate of 20/s through an ear-inserted transducer, are non-filtered alternating clicks of 80 or 90 dB HL (the intensity being kept constant during a monitoring session). BAEPs are analog filtered with a bandpass of 150–1300 Hz. Then an adaptive digital filter based on Wiener’s filtering theory is calculated automatically and readapted by the computer for each new BAEP. This adaptive filter usually obtains interpretable traces after 200 to 400 stimulations (fig 1), thus reducing the recording time to 10–20 s for each individual BAEP.

At the beginning of the monitoring session 5 to 10 standard BAEPs were recorded to check the reliability and reproducibility of the responses. Waves I, III and V were identified by the examiner on these first records and their latencies, calculated by means of a cursor, served as reference values for subsequent automatic peak detection. The rate of BAEP...
recording was usually of 1 new EP every 3–5 minutes, but could be modified at any moment during monitoring. The last 10 responses obtained, as well as the trend curves of latencies corresponding to the last 24 hours of monitoring, were permanently displayed on the computer screen at the patient’s bedside (fig 1). BAEP recording, adaptive digital filtering, automatic peak detection, disk storage and screen display were usually performed in less than 1 minute. Intracranial pressure (ICP) values were also digitised and displayed on the screen along with BAEPs.

b) Patients
Fifty seven patients, all of them deeply comatose (Glasgow Coma Scale ≤ 6 in 54 cases, < 8 in the others) were monitored during periods ranging from 5 hours to 13 days. The mean (SD) duration of monitoring was 98 (50) hours, that is about four days; the mean number of BAEP recordings obtained per patient was 1920 (range: 40–7000). The aetiology of coma was supratentorial in 55 patients (97%), while a primary brainstem lesion could be demonstrated in the remaining two. The origin of coma was traumatic in 35 cases (61%) and vascular in 22 (39%). The patients’ mean (SD) age was 35 (16) (range 6–64). As expected, the traumatic group was significantly younger, mean (SD) years = 27 (13) than the vascular group, 46 (14) years. Metabolic or anoxic comas were not included in the study; our conclusions therefore only concern coma of traumatic or vascular origin.

Clinical outcome was evaluated at three months. Death occurred in 29 cases (50.8%) as a direct result of the brain lesions. Three patients (5.3%) developed a persistent vegetative state (PVS), of whom one died two years later. Finally, 4 patients (7%) died from non neurological causes apparently unrelated to the aetiology of coma (generalised sepsis in 2 and cardiac failure in the other 2). The rate of unfavourable outcome (death or PVS) was significantly greater within the stroke group than in head injured subjects (77% vs 53% respectively, p < 0.05).

In 53 patients (93% of the series) ICP and arterial pressure were also continuously monitored. ICP values were obtained by means of an epidural sensor13 and stored on floppy disks along with the neurophysiological parameters. Cerebral perfusion pressure (CPP) was computed as the difference between mean arterial and intracranial pressures (MAP-ICP). Due to technical mishaps the ICP values were considered unreliable in 2 cases by the intensive care practitioner (FA); consequently correlations with BAEP could only be attempted in 51 patients.

c) Normative data and abnormality criteria

Single BAEPs
Normative data were obtained from 40 control subjects with normal hearing.14 For each response the I-V interpeak latency (IPL), the V/I amplitude ratio and the amplitude of waves III and V were assessed, with normal limits set at the mean value ± 3 SD of our control group. Due to possible injury to the peripheral auditory system in traumatic coma, the absence of
any reproducible potential was considered as significant only if preceded by a progressive loss of components verified during monitoring.

The prognostic implications of single BAEPs were studied in two different sets of responses, recorded respectively during the first 48 hours of coma and at the beginning of the second week (day 6–7 of coma). These two sets of responses were respectively labelled "early" and "late" BAEPs; they were investigated separately to assess whether the moment at which recordings were made could influence their prognostic value. In a subset of 15 patients data from both, early and late recording sessions were available. In them we could investigate the prognostic relevance of BAEP changes occurring over the first week.

**Dynamic study of BAEPs**

Normative data for the dynamic study of BAEPs were derived from 10 normal hearing subjects in whom auditory responses were recorded during 8 to 10 consecutive hours at a rate of 1 BAEP every 10 minutes. EEG and rectal temperature were also monitored in these subjects to assess the physiological variations of responses related to body temperature and vigilance status. A total of 590 BAEPs were obtained [mean (SD) 58 (10) per subject], including recordings during physiological sleep in all subjects. BAEP latencies during monitoring were best characterised by their mean (SD) value, this latter providing a criterion of latency stability. In our control subjects SD was 0-15 ms for wave V latency, and 0-1 ms for I-V IPL.

Transient modifications of the I-V interval detected during monitoring were considered significant when exceeding 2-5 SD, on the condition that at least 10 consecutive BAEPs were available during the increase and decrease of latency, so that a regression line could be calculated for the successive latency values in each phase.

BAEP modifications related to body temperature changes or observed under blood levels of anaesthetic drugs known to alter BAEPs were excluded, after previous studies in which the effects of such non-pathological factors on BAEP monitoring were systematically assessed.

**ICP and CPP changes**

Acute pressure modifications were considered significant on the two conditions: 1) that ICP raised beyond 40 mmHg or CPP decreased below this same level, and 2) that the episode lasted a minimum of 10 minutes. On these criteria we recorded pressure abnormalities in 26 patients of the series (42-3%); BAEP latency evolution during these episodes could be reliably studied in all but four of them, in whom the brainstem responses were either abolished (one patient) or altered by high doses of drugs (three cases) at the moment that ICP rise took place.

### Results

#### A "Static" study of BAEPs during coma.

**Prognostic implications**

1) **Initial BAEP and life outcome**

After excluding patients under continuous infusion of anaesthetic agents which altered significantly the responses, the relationship between "early" BAEPs (obtained during the first 48 hours of coma) and clinical outcome was studied in 37 patients.

Twenty seven patients had initially normal BAEPs and 14 (51-8%) recovered full consciousness, while in 13 cases (48-1%) the evolution was towards death. Within this group of normal BAEPs, neither the mean age, nor the I-V interval at monitoring onset could discriminate between patients with good and bad outcome (table 1).

BAEPs were initially abnormal in 10 patients. Nine of these had supratentorial hemispheric lesions, who all died. The remaining patient had an isolated haemorrhagic brainstem contusion that could explain BAEP changes, and recovered from coma.

2) **"Late" BAEPs and outcome**

"Late" BAEPs (obtained during the second week of coma) were recorded in 30 patients. Four of them had abnormal or absent responses, all of who died. Of 26 patients with

#### Table 1 Comparison (unpaired t test) of several variables within the groups of "early" (n = 27) and "late" (n = 26) normal BAEPs. In patients with "early" normal responses (left half of the table) neither the age nor the interpeak latencies (I-V IPL) could discriminate between survivors and non-survivors. Conversely, in patients with "late" normal BAEPs the I-V interval was significantly longer in those who subsequently died. Values in the two lower rows are mean (SD). Age in years, I-V IPL in milliseconds.

<table>
<thead>
<tr>
<th></th>
<th>&quot;Early&quot; Normal BAEPs (first 24 hour) (n = 27)</th>
<th>&quot;Late&quot; Normal BAEPs (6th-7th day) (n = 26)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Survival</td>
<td>Death</td>
</tr>
<tr>
<td></td>
<td>N (%)</td>
<td></td>
</tr>
<tr>
<td>N (%)</td>
<td>13 (48-1)</td>
<td>14 (51-1)</td>
</tr>
<tr>
<td>Age</td>
<td>33 (17)</td>
<td>30 (17)</td>
</tr>
<tr>
<td>I-V IPL</td>
<td>3-97 (0-3)</td>
<td>4-03 (0-3)</td>
</tr>
</tbody>
</table>

Figure 2 Temporal evolution of the I-V interval (IPL) in 15 patients continuously monitored during the first 6-10 days of coma. Values from patients who survived (open circles) and from those who subsequently died (filled circles) are presented separately. In the survivors group the I-V IPL remained stable between the first 48 hours and the seventh day of coma (open circles), while this measure increased significantly in the group of patients who subsequently died (filled circles, the cross means p < 0.05 on paired t test). Clinical outcome was assessed at three months. Note that, despite the changes operated between the two recording sessions, the interpeak latencies remained within normal limits (mean ± 3 SD) in both groups.
normal "late" BAEPs 13 survived and 13 died (4 of them from presumed non-neurological causes). Thus a normal BAEP recorded beyond the first days of coma did not warrant a good outcome in our patients. However, the I-V interval at one week's evolution was significantly shorter in the patients who survived than in those who died (unpaired t test, p < 0.05, see table 1), although it remained within normal limits in both cases. The mean age was comparable in the first and in the second group.

When data from the "early" (first 48 hours) and "late" (second week) recording sessions were compared no significant modification of the I-V IPL was observed in the survivors, whereas patients who subsequently died showed a significant increase of this parameter (paired t test p < 0.05, fig 2).

Brain death occurred in only 2 of the 9 patients (22%) with either no visible lesions on CT scans, or only extracerebral lesions such as epi- or subdural haematomas, and who showed normal "late" BAEPs. Conversely, death occurred in 11 of 17 patients (65%) with normal BAEPs on the second week of coma and intracerebral haemorrhagic lesions (cerebral haemorrhages or haemorrhagic infarcts)

B) Dynamic study of BAEP changes during coma
1) Transient BAEP changes
Applying the criteria specified in the methods section, 25 episodes of transient, reversible latency changes of the I-V IPL were observed in 18 patients of our series (figs 3 and 4). Despite such latency increase the I-V IPL remained within normal standard limits (<4.7 ms) in all except one patient. The duration of IPL variation episodes ranged from two to 12 hours. Only interpeak latencies were taken into account since amplitudes were in our experience too susceptible to uncontrolled sources of variations to be fully reliable.

The overall mortality in this group was significantly higher than that observed in the patients who did not present transient abnormalities of the I-V IPL (14/18 (78%) vs 16/33 (48.5%), p < 0.05). This difference was still increased if only patients with normal BAEPs at monitoring onset were considered, in which case the mortality associated with transient IPL increases was of 76%, in contrast with 37% for all other patients. The mean (SD) age of patients in both groups was not significantly different, [34.6 (17) vs 35.6 (16) years] and could not account for the difference in outcome.

2) Acute ICP/CPP modifications associated to BAEP changes
As shown in table 2, patients with acute increases of the I-V interval were more likely to have ICP/CPP modifications during monitoring than those without BAEP changes. Conversely, the existence of significant episodes of pressure modification was not associated with a higher proportion of BAEP abnormalities.

Figure 3  Long-lasting, transient latency increase during coma monitoring. The left part of the figure shows the trend curves of BAEP latencies (three upper graphs) and of intracranial pressure values (bottom line). The onset of the episode of BAEP latency increase (A) coincided with an acute rise of ICP (arrow on bottom trace) but lasted several hours longer. Letters A, B and C mark corresponding periods of monitoring in trend curves (left) and BAEP traces (right). Thus, traces on the right are representative BAEPs obtained at the onset (A), the moment of maximal latency increase (B) and the end of the episode (C). Each illustrated trace corresponds to the average of 5 consecutive BAEPs (1000 stim, 10 minutes of monitoring). Note that, in spite of the significant effect seen on trend curves the actual increase of wave V latency on BAEP traces was very small. After the end of the episode a slow latency increase developed in the absence of new ICP changes, but was enhanced when a new ICP rise appeared at 6 am. BAEP monitoring was discontinued at 10 am, when BAEPs were still present, the patient died six hours later.
Only in half of the patients were pressure changes followed by acute BAEP latency increase; in the others, BAEP remained stable during and after the episode of ICP/CPP alterations (Table 2).

Reversible increases of the I-V interval were time-locked to the ICP/CPP changes in 11 patients, of whom 10 (90.9%) had an unfavourable outcome towards cerebral death. In 11 other patients BAEP latencies remained unchanged in spite of acute pressure modifications. Nine of them (82%) survived. From the two patients who died one had very abnormal BAEPs from the beginning of monitoring and the remaining died because of an intercurrent sepsis. In the remaining four patients BAEP evolution during acute ICP changes could not be assessed, either due to the high doses of sedative drugs which prevented BAEP interpretation (three cases), or because of BAEPs being already abolished at the time of ICP rise (one case). Thus the association of acute ICP/CPP modification plus I-V increase was followed by neurological death in 91% of cases, whereas mortality was only 18% when acute ICP/CPP changes were not accompanied by transient BAEP latency increase. The difference is highly significant (p < 0.001).

Table 2 Distribution of the 57 patients monitored according to the existence of a) transient modifications of intracranial or cerebral perfusion pressures, and b) transient increases of the I-V interpeak latency. The existence of latency increase was very frequently (11/15) accompanied by pressure changes; however, pressure modifications were only followed by a BAEP latency increase in 50% of cases. A discussion in the text, an unfavourable outcome was associated to the concomitant occurrence of BAEP and pressure changes.

<table>
<thead>
<tr>
<th>I-V IPL increase</th>
<th>Yes</th>
<th>No</th>
<th>ND*</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICP non controlled</td>
<td>11</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>BAEP absent or unreliable due to high levels of lidocaine/ thiopental</td>
<td>11</td>
<td>20</td>
<td>2</td>
</tr>
<tr>
<td>ND* = Not done.</td>
<td>4</td>
<td>1</td>
<td>1</td>
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**Discussion**

A) Prognostic implications of single BAEP recordings at different moments of the evolution

Gross abnormalities or abolition of BAEPs in a comatose patient carry an ominous prognosis, provided that lesions to the peripheral auditory system have been ruled out. The poor prognosis associated with this pattern during coma is illustrated by the 90% mortality observed in our group of patients with abnormal BAEPs at monitoring onset.

The central latency of BAEPs has been reported as being almost constantly normal when recorded during the first six hours of traumatic coma. This suggests that most abnormalities encountered later are secondary to hemisphere lesions, and reflect an ongoing process of rostrocaudal deterioration of the brainstem. Discrete brainstem lesions may, however, also induce severe BAEP abnormalities if they involve the central auditory pathways. BAEP abnormalities in those cases reflect the primary brainstem lesion, and not the rostrocaudal deterioration of brainstem function, and the ensuing outcome is not necessarily bad. This was the case of one of our patients with severely abnormal BAEPs resulting from a small pontine haemorrhage, without evidence of hemispheric lesion, who recovered from coma some weeks later.

Contrary to what happens with abnormal brainstem responses, normal BAEPs during the first days of coma do not bring much relevant information on the final outcome. In published series the average incidence of unfavourable evolution (severe disabilities, PVS or death) in patients with bilaterally normal BAEPs during the acute phase of traumatic coma is about 30%, but ranges from 0 to more than 50%. In our study 46% of patients with normal “early” BAEPs did unfavourably, a figure comparable to other series dealing with patients having low Glasgow Coma Scores.

Some studies have suggested that a favourable vital outcome might be predicted by normal BAEPs recorded beyond the first days of coma. The reason for this is probably that death after traumatic or vascular coma mainly occurs during the first week, and especially during the first 48 hours after the injury.
BAEP and ICP monitoring in coma

We addressed this question by investigating 26 patients of our series who had normal "late" BAEPs recorded during the second week of coma, 15 of whom had also been studied within the first 48 hours.

When the group of 26 patients is considered as a whole, "late" normal BAEPs do not prove of greater prognostic value than "early" normal responses, since mortality was still 50% (13 patients). However, the nature of the lesion might also influence the prognostic value of normal "late" BAEPs, since death occurred in only two of nine cases (22%) without focal injury on CT scans, or with extracerebral lesions not associated to intraparenchymal involvement, whereas brain death ensued in 11 of 17 patients (65%) who had intracerebral haemorrhagic lesions, in spite of persisting normal BAEP's during the second week. Thus only in a small subset of patients may normal "late" BAEPs be associated with a good vital outcome. This subgroup did not include patients with intracerebral haemorrhages or haemorrhagic infarcts, which is not surprising since such lesions have the highest potentiality for late neurological deterioration. As a general rule, the smaller the probability of a secondary insult (such as, late ICP rise, rebleeding) the better the prognostic value of a "late" normal BAEP.

The expected outcome may be optimised by assessing the dynamics of brainstem responses, rather than their statistical normality at any time of coma evolution. One of the main reasons counteracting to the lack of prognostic power of normal BAEPs is that "normal" waveforms are in many cases progressively deteriorating responses that will remain within normal limits for a long time. This point is illustrated by fig 2, which shows central latency changes in our patients between the first and second weeks of coma. Although latencies increased significantly in the group of patients who subsequently died, all values remained within statistical normal limits in both recording sessions, this stresses the limitations of statistical "normality" when applied to comatose patients.

B) Transient BAEP changes during coma

Transient BAEP changes lasting from minutes to hours are frequent in comatose patients. Many of these are non-pathological in origin and mainly related to temperature or drug levels modifications; however, detection of clinically relevant episodes is crucial, for they may represent the earliest signs of brainstem dysfunction, especially if associated with critical alterations of other markers such as ICP or CPP.

From our previous work on comatose patients and normal subjects we devised criteria to distinguish changes related to CNS dysfunction from those of non-pathological origin. In this series of patients latency changes fulfilling our quite restrictive criteria did not necessarily make BAEP exceed statistical norms, as established in a control population. In most patients latencies remained within normal limits during the whole episode, and would have not been considered as abnormal had a single BAEP been obtained at that moment. We have previously demonstrated that in patients with irreversible changes progressing up to brain death BAEP abolition mostly occurred before the I-V IPL had exceeded normal values. This emphasises the fact that during monitoring relative changes from baseline are much more important than comparisons to normative data banks.

In our patients, the significance of transient latency increases was best assessed when studied in correlation with ICP and CPP. Only 50% of acute ICP/CPP abnormalities were accompanied by an increase of the I-V interval, thus suggesting that not all "significant" pressure changes entail a functional suffering of the brainstem, and that the mere monitoring of ICP values may not always predict their actual incidence on brainstem function. This is consistent with the results of Kawahara et al, who reported that the range of ICP values associated to wave V loss during BAEP monitoring were extremely wide, and could range from 30 to 150 mmHg. In our patients it was clear that for the same category of pressure changes, the ensuing clinical evolution differed drastically depending on whether a concomitant BAEP modification appeared or not. When ICP changes were accompanied by transient I-V increases the subsequent clinical evolution was almost constantly bad, with more than 90% progression to brain death in the following hours or days, whereas outcome was significantly better (18% brain death or PVS) in patients whose episodes of pressure change did not entail BAEP modifications.

Transient displacement of the brainstem, either laterally due to uncial herniation or caudally during central herniation, may entail the development of secondary intraparenchymal haemorrhages due to stretching and tearing of perforant branches of the basilar artery. In dogs submitted to transient ICP rises followed by reperfusion of the perforant arteries, mesencephalic haemorrhages developed due to blood extravasation at the moment of reperfusion, which are located lesions of brainstem arteries due to the preceding stretching. We suggest that in our patients, transient brainstem distortion secondary to acute ICP rise, and manifested by BAEP changes, might have set up an ongoing and self-perpetuated process, with progressive brainstem deterioration culminating in brain death. This would be consistent with the slow progressive latency increase that followed transient BAEP episodes in several patients (fig 3), and which ultimately lead to brain death in the absence of further ICP/CPP modifications.

At the time our recordings were obtained the detection of transient BAEP changes did not entail a modification of patient's management; therefore we cannot ascertain whether these acute changes are the first signs of a potentially reversible process, or conversely herald an irreversible brainstem deterioration. What seems certain is that BAEP monitoring provides important information on whether a particular episode of ICP rise is affecting
brainstem function or not. The association of acute ICP rises to BAEP modifications, even if both are transient and totally reversible, is a reliable sign of alarm. In those cases a significant brainstem suffering may be assumed even if pressure modifications do not reach the common clinical assumptions of "critical threshold" values as BAEP latency changes in that context seem more reliable a warning than absolute intracranial pressure levels.

From this study it may also be concluded that BAEP stability is the only desirable finding during coma monitoring, since any significant modification, even transient, of BAEPs implicates a very poor outcome. It seems reasonable, therefore, to suggest that, in addition to BAEPs, the monitoring of hemispheric function with cortical EPs must be considered to detect the signs preceding irreversible brainstem damage. Technically this can be implemented either by simultaneous monitoring of short- and middle-latency responses (auditory or somatosensory), or by starting monitoring with cortical EPs, and shifting to (or adding) brainstem recordings when the cortical components become altered or disappear.