Brain Response to One’s Own Name in Vegetative State, Minimally Conscious State, and Locked-in Syndrome

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**Background:** A major challenge in the management of severely brain-injured patients with altered states of consciousness is to estimate their residual perception of the environment.

**Objective:** To investigate the integrity of detection of one’s own name in patients in a behaviorally well-documented vegetative state (VS), patients in a minimally conscious state (MCS), and patients with locked-in syndrome.

**Design:** We recorded the auditory evoked potentials to the patient’s own name and to 7 other equiprobable first names in 15 brain-damaged patients.

**Results:** A P3 component was observed in response to the patient’s name in all patients with locked-in syndrome, in all MCS patients, and in 3 of 5 patients in a VS. P3 latency was significantly (P<.05) delayed for MCS and VS patients compared with healthy volunteers.

**Conclusions:** These results suggest that partially preserved semantic processing could be observed in non-communicative brain-damaged patients, notably for the detection of salient stimuli, such as the subject’s own name. This function seems delayed in MCS and (if present) in VS patients. More important, a P3 response does not necessarily reflect conscious perception and cannot be used to differentiate VS from MCS patients.

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words (presented in a sequence of multiple occurrences of a single word), notably when they are pertinent, such as for the subject’s own first name. However, the presence of a P3 wave in such paradigms could reflect either the recognition of the target’s intrinsic meaning or the detection of its acoustic salience (ie, the fact of being rare relative to a monotonous series) because the P3 amplitude is sensitive to task relevance and stimulus probability. For a review, see the study by Polich and Kok. To avoid this ambiguity, it has been proposed to remove the physical rarity of the target stimuli by using as sensory input a series of equiprobable first names among which is included the subject’s own name (SON). In these conditions, P3 presence becomes a valid neurophysiologic correlate of word semantic categorization.

Our current study explores the integrity of SON discrimination (independently of target occurrence probability) in severely brain-damaged coma survivors (those in a VS, those in a minimally conscious state [MCS], and those with locked-in syndrome [LIS]). Furthermore, we aimed to objectively assess, individually and at the bedside, the possible preservation of residual linguistic processing differentiating unconscious VS from MCS patients by an objective electrophysiological processing differentiating unconscious VS from MCS patients by an objective electrophysiological measurement.

METHODS

SUBJECTS

This study was prospectively performed in 18 right-handed severely brain-damaged patients classified according to internationally established criteria as being (1) in a VS,10 (2) in an MCS,11 or (3) affected by LIS.12 Only patients studied during awake periods, free of centrally acting drugs and without diagnostic ambiguity, were included for further analysis. Each data set composed cognitive ERP measurements and standardized clinical assessments of consciousness.

Fifteen patients (mean±SD age, 54.9±17.2 years) were included for further analysis (3 patients were excluded because of technical problems): 5 were in VS (4 nontraumatic and 1 traumatic) (mean±SD age, 51.8±13.0 years), 6 were in MCS (3 nontraumatic and 3 traumatic) (mean±SD age, 58.5±19.5 years), and 4 were affected by LIS (all traumatic) (mean±SD age, 53.3±21.9 years). (Table 1). All patients were right-handed, as evaluated by heteroromanames. None had a history of impaired auditory acuity. Somatosensory evoked potentials obtained by stimulation of the median nerve showed the presence of primary somatosensory cortex potentials (N20) in all patients. Five age-matched right-handed (Edinburgh inventory) healthy volunteers (2 women and 3 men; mean±SD age, 54.6±11.3 years) participated in the experiment. None of them had a history of audiological or neurological disease. The experiment was conducted in agreement with the guidelines of the Declaration of Helsinki and was approved by the Ethics Committee of the Faculty of Medicine of the University of Liège. Written/eye-coded informed consent was obtained from all subjects or LIS patients or from a member of the patient’s family.

STANDARDIZED BEHAVIORAL EVALUATION

Before ERP measurements, an experienced neuropsychologist (C.S.) performed behavioral testing by the Glasgow-Liège Scale with a quantified analysis of brainstem reflexes: fronto-orbicular, vertical oculocephalic, pupillary, horizontal oculocephalic, and oculocardiac. The Glasgow-Liège Scale is calculated as the sum of eye opening, motor response, verbal response, and brainstem reflex subscores, and is scored from 3 (worst) to 20 (best). The JFK Coma Recovery Scale—Revised is a recently validated behavioral scale that explicitly incorporates the diagnostic criteria for VS10 and MCS11 into its administration and scoring scheme, and is unique in allowing the derivation of a diagnostic directly from the examination findings. It includes auditory, visual, motor, oromotor/verbal, communication, and arousal subscales, and ranges from 0 (worst) to 23 (best). Following ERP recordings, neurobehavioral evaluation continued for VS and MCS patients twice a month to increase diagnostic certainty.

STIMULATIONS

We elaborated 6 sequences of 80 stimulations containing 8 first names, 1 of them always being the SON. Only this name changed for every subject, the 7 other first names (OFNs) being the same for all participants. The OFNs were selected from a previous study as a series of first names of similar high frequency of use in the French language. For each sequence, each name was presented 10 times in random order, thus making a complete series of 80 equiprobable first names (probability, 12.5% for each name), with an interstimulus interval varying between 1000 and 1400 milliseconds. After each recording session, the subject or the subject’s family was asked whether 1 of the other names had a particular emotional importance (ie, they corresponded to names of close relatives). If it was the case, the name was excluded from the ERP analysis. All first names were di-syllabic, were recorded by the same neutral male voice (F.P.), and were digitized and replayed binaurally at a 90-dB sound pressure level maximal intensity.

ERP ACQUISITION

For patients, data were acquired at their bedside. Preceding each electroencephalographic recording, the behavioral status was evaluated by the JFK Coma Recovery Scale—Revised and the Glasgow-Liège Scale. Control subjects were studied while laying in bed with minimal ambient noise. Electrodes and mini-earphones were put in place. The electroencephalographic signals from 3 electrodes, Fz, Cz, and Pz (placed according to the International Ten-Twenty System), referenced to the nose; the electro-oculogram, from 2 electrodes diagonally above and below the right eye; and the electromyogram, from 2 electrodes on the chin were amplified (×150 000) and sampled at 300 Hz by an acquisition system (NuAmps; NeuroSoft, Sterling, Va) with an analog bandpass of 0.1 to 70 Hz (except for the electromyogram where the bandpass was 10-100 Hz). A ground electrode was placed near Fz, and impedances were kept below 5 kΩ. The subjects heard, eyes closed, 6 series of 80 equiprobable first names, without any specific task (ie, passive condition).

ERP ANALYSIS

Event-related potentials were averaged according to the type of first name (SON vs OFN) and the electrode position (Fz vs Cz vs Pz). Before averaging, single epochs with an amplitude of 50 μV or higher or those containing eye movements or electromyographic artifacts were excluded from averaging. A different number of trials between SON and OFN may
bias our obtained results because of signal-noise ratio differences. To address this concern, separate OFN averages were constituted, each containing a similar number of trials as the SON average. Comparisons between separate OFN and SON averages confirmed results obtained with grand averages of OFN data, validating our reported results. For illustrative purposes, grand-averaged ERPs were constructed for all control subjects and for each clinical entity (VS, MCS, and LIS).

Table 1. Demographic, Clinical, and ERP Data

<table>
<thead>
<tr>
<th>Patient No./Sex/Age, y</th>
<th>Clinical Diagnosis</th>
<th>Cause</th>
<th>GCS Score on Admission*</th>
<th>MRI Findings</th>
<th>EEG Findings (Background Activity)</th>
<th>Time Spent Before Examination</th>
<th>JFK CRS-R Score†</th>
<th>GLS Score‡</th>
<th>GOS Score (1 y After ERP)§</th>
<th>P3 Present</th>
</tr>
</thead>
<tbody>
<tr>
<td>1/M/29</td>
<td>VS</td>
<td>Trauma</td>
<td>3</td>
<td>Mesencephalic lesion and right frontal and left cerebellar hemorrhages</td>
<td>Disorganized δ</td>
<td>14 d</td>
<td>3</td>
<td>11</td>
<td>2</td>
<td>Yes</td>
</tr>
<tr>
<td>2/M/55</td>
<td>VS</td>
<td>Cardiorespiratory arrest</td>
<td>5</td>
<td>Perventricular white matter lesions</td>
<td>Diffuse δ</td>
<td>3 mo</td>
<td>6</td>
<td>12</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>3/M/57</td>
<td>VS</td>
<td>Viral encephalitis</td>
<td>6</td>
<td>Normal</td>
<td>Diffuse δ and α</td>
<td>19 d</td>
<td>5</td>
<td>12</td>
<td>2</td>
<td>No</td>
</tr>
<tr>
<td>4/M/62</td>
<td>VS</td>
<td>Cardiorespiratory arrest</td>
<td>6</td>
<td>Bilateral</td>
<td>Perventricular lesions, hydrocephalus, and diffuse cortical atrophy</td>
<td>Diffuse δ</td>
<td>8.5 mo</td>
<td>6</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>5/F/56</td>
<td>VS</td>
<td>Intracerebral hemorrhage</td>
<td>5</td>
<td>Left frontal and intraventricular hemorrhage</td>
<td>Disorganized δ and left predominant δ</td>
<td>1.5 mo</td>
<td>7</td>
<td>13</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>6/F/83</td>
<td>MCS</td>
<td>Intracerebral hemorrhage</td>
<td>9</td>
<td>Bilateral fronto-temporal and perilesional edema</td>
<td>Low-voltage δ</td>
<td>18 d</td>
<td>12</td>
<td>14</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>7/M/72</td>
<td>MCS</td>
<td>Respiratory insufficiency</td>
<td>9</td>
<td>Diffuse perventricular and right caudate lesions</td>
<td>Disorganized δ</td>
<td>2 mo</td>
<td>10</td>
<td>15</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>8/M/34</td>
<td>MCS</td>
<td>Trauma</td>
<td>4</td>
<td>Bilateral contusion, pontine lesion, and normotensive hydrocephalus</td>
<td>Disorganized δ</td>
<td>10 mo</td>
<td>11</td>
<td>16</td>
<td>3</td>
<td>Yes</td>
</tr>
<tr>
<td>9/M/49</td>
<td>MCS</td>
<td>Trauma</td>
<td>5</td>
<td>Left temporoparietal extradural hematoma and right temporal and fronto-contusions</td>
<td>Disorganized δ and left predominant δ</td>
<td>20 d</td>
<td>15</td>
<td>15</td>
<td>5</td>
<td>Yes</td>
</tr>
<tr>
<td>10/M/71</td>
<td>MCS</td>
<td>Trauma</td>
<td>7</td>
<td>Right temporoparietal hematoma and perilesional edema</td>
<td>Disorganized δ and diffuse δ</td>
<td>13 d</td>
<td>12</td>
<td>15</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>11/M/42</td>
<td>MCS</td>
<td>Intracerebral hemorrhage</td>
<td>3</td>
<td>Left frontal hemorrhage (drained) and multifocal bilateral lesions</td>
<td>Disorganized δ</td>
<td>2.5 mo</td>
<td>16</td>
<td>17</td>
<td>1</td>
<td>Yes</td>
</tr>
<tr>
<td>12/M/77</td>
<td>LIS</td>
<td>Basilar artery thrombosis</td>
<td>3</td>
<td>Pontomesencephalic lesion</td>
<td>Disorganized δ</td>
<td>1 mo</td>
<td>16</td>
<td>11</td>
<td>3</td>
<td>Yes</td>
</tr>
<tr>
<td>13/F/24</td>
<td>LIS</td>
<td>Basilar artery thrombosis</td>
<td>6</td>
<td>Ventral pontine lesion</td>
<td>α</td>
<td>5 y</td>
<td>16</td>
<td>12</td>
<td>3</td>
<td>Yes</td>
</tr>
<tr>
<td>14/M/57</td>
<td>LIS</td>
<td>Basilar artery thrombosis</td>
<td>6</td>
<td>Pontomesencephalic lesion</td>
<td>α</td>
<td>7 y</td>
<td>16</td>
<td>12</td>
<td>3</td>
<td>Yes</td>
</tr>
<tr>
<td>15/M/55</td>
<td>LIS</td>
<td>Traumatic basilar artery dissection</td>
<td>4</td>
<td>Pontomesencephalic lesion</td>
<td>α and moderate diffuse δ</td>
<td>3 y</td>
<td>16</td>
<td>12</td>
<td>3</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Abbreviations: CRS-R, Coma Recovery Scale–Revised; EEG, electroencephalogruphic; ERP, event-related potential; GCS, Glasgow Coma Scale; GLS, Glasgow-Liège Scale; GOS, Glasgow Outcome Scale; LIS, locked-in syndrome; MCS, minimally conscious state; MRI, magnetic resonance imaging; VS, vegetative state.

*The total possible was 15.
†The total possible was 23.
‡The total possible was 20.
§The total possible was 5 (1 indicates dead; 2, VS; 3, severely disabled; 4, moderately disabled; and 5, good recovery).13
SON for certain patients), we chose the maximum amplitude (and its associated latency) in a temporal window predefined on grand-averaged ERPs. Amplitude values were tested with a multivariate analysis of variance (MANOVA) with repeated measures on component (N1 vs P2 vs N2 vs P3), name (SON vs OFN), and electrode position (Fz vs Cz vs Pz). Latency values were tested with a MANOVA with repeated measures on component (N1 vs P2 vs N2 vs P3) and name (SON vs OFN). The independent variable was the group (control, MCS, VS, and LIS). The MANOVAs were subjected to a Greenhouse-Geisser conservative df correction. A Tukey comparison with a threshold at P<.05, and corrected for multiple comparisons, was performed when significant interactions emerged on MANOVA.

In addition to the visual inspection and interpretation of the data, a statistical analysis tested in each subject the significance of the P300 response to the SON compared with the OFN. Adopting a similar approach as Marchand et al., individual waveforms were analyzed on a point-by-point basis using serial t scores, which take into account the variance of the individual electroencephalographic trials composing the grand-averaged ERP. t Scores were computed for all subjects in a temporal window of 50 milliseconds around the peak latency of the P300. Results were considered significant at P<.05.

### RESULTS

In healthy controls, the SON evoked the classic N1, P2, N2, and P3 components at about 150, 230, 305, and 460 milliseconds, respectively (Table 2 and Figure 1). The ERP latencies were delayed compared with those reported in the literature because the population of our study was older; this effect being previously well documented.

Among the patients, in all but 2 cases (patients 3 and 4, both in a VS), well-defined ERP components were obtained in individual averages (Figure 2). Notably, a P3 component was clearly observed after the SON in all patients affected by LIS and in an MCS, and in 3 of 5 patients in a VS (Figures 1 and 2). The individual statistical P300 analysis showed significant t scores (P<.05) for all subjects, except for 2 VS patients (patients 3 and 4) and 1 LIS patient (patient 13). The LIS patient failed to show a significant P300 (P=.06), probably owing to the recording having many artifacts and the ensuing insufficient statistical power (because of uncontrollable microwave muscle contractions, only 13 valid trials could be obtained for the SON in this case).

The MANOVA did not show any significant (P=.65) group effect on ERP amplitudes. In contrast, the analysis demonstrated that the interaction between component and name had a significant effect on ERP amplitudes (F3,98 = 26.53, P<.001). Post-hoc analyses revealed, as expected, that P3 amplitude was significantly higher in response to SON than to OFNs for all groups (P<.05) (Table 3). No significant differences (P>.05) were observed for the other components.

With respect to latencies, the MANOVA showed a significant group × component interaction (F9,48 = 3.5, P=.01). Post-hoc analyses revealed that P3 latency was significantly delayed for the VS group compared with the LIS group and controls, and for the MCS group compared with the controls (Table 2). No significant differences (P>.05) were observed for the other components.

### COMMENT

It is not surprising to observe a differential P3 wave in LIS patients because it can be expected that their cognitive functions, and notably their linguistic comprehension, remain preserved, even if their bedside cognitive neuropsychological testing remains challenging.

For MCS patients, our results extend those of Boly et al. and Schiff et al., who suggested that the auditory system of these patients was relatively well preserved in response to passive tones and language stimulation, respectively. The emergence of a P3 wave to the SON (compared with OFNs) suggests that MCS patients are able to detect salient words. However, the significant difference in P3 latency compared with controls shows that this word processing is delayed in the MCS group.

The many behaviorally well-documented VS patients emitting a differential P3 was somehow unexpected. Previous studies have reported fewer unconscious patients emitting a P3 to deviant (and rare) tones. Yingling et al., Harris and Hall, Gott et al., and Mut-schler et al. were able to record P3 components in 22% to 30% of comatose patients (2 of 9, 2 of 8, 6 of 20, and 6 of 20 patients, respectively). Similarly, Glass et al. have recorded a P3 response using an ordinary oddball paradigm in 38% (3 of 8) of patients thought to be in a VS. The presence of a P3 response in 3 of 5 of our VS patients might be because of the quality of the implicitly targeted stimuli (ie, the SON), which was highly salient because of its obvious emotional dimension and its familiarity. Accordingly, Signorino et al. have shown that emotional stimuli increase the chances of obtaining a P3 response in comatose patients. These researchers used a conventional oddball paradigm and an oddball paradigm in which the stimuli were coupled to emotional verbal stimuli (ie, a short phrase spoken by a member of

### Table 2. Data for N1, P2, N2, and P3 Latencies in Response to the Subject’s Own Name for Each Clinical Entity

<table>
<thead>
<tr>
<th>Group</th>
<th>N1</th>
<th>P2</th>
<th>N2</th>
<th>P3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Those in a VS</td>
<td>191.2 ± 33.7</td>
<td>332.8 ± 30.2</td>
<td>475.6 ± 26.5</td>
<td>762.4 ± 35.2</td>
</tr>
<tr>
<td>Those in an MCS</td>
<td>178.4 ± 14.4</td>
<td>286.4 ± 29.6</td>
<td>423.6 ± 36.2</td>
<td>711.2 ± 57.8†</td>
</tr>
<tr>
<td>Those with LIS</td>
<td>151.0 ± 9.7</td>
<td>202.5 ± 9.1</td>
<td>296.5 ± 16.7</td>
<td>531.0 ± 52.7</td>
</tr>
<tr>
<td>Controls</td>
<td>148.4 ± 5.0</td>
<td>232.4 ± 19.3</td>
<td>306.8 ± 27.0</td>
<td>460.6 ± 33.7</td>
</tr>
</tbody>
</table>

Abbreviations: See Table 1.

*Data are given as mean ± SE milliseconds.

†P<.05 compared with controls.
the family or the patient’s name) and obtained a P3 in 38% (6 of 16) of comatose patients in the first condition and in 56% (9 of 16) of comatose patients in the second condition. Our study further suggests that salient stimuli, such as the SON, can induce linguistic processing in some clinically well-documented VS patients, although delayed compared with age-matched controls. However, the absence of significant differences between the VS and MCS groups indicates that the P3 response cannot be used to differentiate VS from MCS or LIS patients. These data emphasize the importance of complementary (nonbehavioral) investigations in the assessment of residual cognition in noncommunicative brain-damaged patients. Patients in a VS are not “apallic”34 or in “neocortical death.”35 The VS is a more heterogeneous clinical entity than previously thought, and VS patients may show preserved islands of functional “pallium” or neocortex. Our findings are also concordant with recent functional neuroimaging work of Schiff et al36 and Owen et al,37 showing that islands of cerebral function may be preserved in some—but not all—VS patients.38

Our results raise several questions. First, which characteristic of the SON stimulus accounts for the brain’s P3 response? Is it familiarity (high frequency of exposure during the entire lifetime) or emotional value? Indeed, the SON is a piece of information that we often process since infancy,39 and it is usually considered to be emotionally charged. Recent work40 suggests that, in healthy subjects, the emotional charge of the SON per

Figure 1. Event-related potentials to the subject’s own name and to other first names in control subjects (n=5) (A), in patients affected by locked-in syndrome (n=4) (B), and in those in a minimally conscious state (n=6) (C) and in a vegetative state (n=5) (D).
se may not be sufficient to grab attention. Thus, it would be useful to evaluate whether a differential P3 component is also recorded when emotional words, rather than the SON, are presented to LIS, MCS, and VS patients.

A second question relates to the conscious perception of the SON. Because the elicitation of a P3 wave is not necessarily concomitant to a phenomenal consciousness (it is also evoked during unconscious subliminal perception),

we would rather limit the interpretation of the observed P3 response in some of our VS patients as an index of partially preserved, albeit restricted, cerebral processing for “automatic” speech comprehension. As suggested by Dehaene and Naccache, phenomenal consciousness is probably the consequence of a coherent activity involving structures distributed throughout the brain. Thus, one way of assessing if noncommunicative patients are aware of external stimuli would be to search for stimulus-induced neural synchronizations using adapted electrophysiological measures.

A last question involves the prognostic significance of a P3 response to presentation of the SON. The few patients who underwent evaluation in the present work

![Figure 2. Individual event-related potentials to the subject’s own name and to other first names in patients in a vegetative state (A-E), in patients in a minimally conscious state (F-K), and in patients affected by locked-in syndrome (L-O). Traces at Cz are represented. A through E represent patients 1 through 5 in Table 1; F through K, patients 6 through 11; and L through O, patients 12 through 15.](http://archneur.jamanetwork.com/pdfaccess.ashx?url=/data/journals/neur/7034/)

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leaves this question open. None of our 5 VS patients subsequently recovered consciousness, and only 1 of the 6 MCS patients showed good recovery 1 year after brain injury. Future investigations in an extended population of patients would help to determine whether this differential P3 response indicates a higher probability of recovery.

In conclusion, to our knowledge, this study is the first to show that a differential P3 component could be recorded in response to the SON, compared with OFNs, in a small but behaviorally well-documented group of MCS and LIS patients (6 of 6 and 4 of 4 patients, respectively) and in some (3 of 5 patients) in a VS. Because we used stimuli that were equiprobable words, the obtained P3 responses can be interpreted as an index of some preserved semantic processing, independently of the probability of occurrence of the stimuli. Our data demonstrate that a P3 response does not necessarily reflect conscious perception and cannot be used to reliably differentiate individuals in a VS from those in an MCS.

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Additional Information: Drs Laureys and Maquet are FNRS research associate and research director, respectively.

Table 3. Data for N1, P2, N2, and P3 Amplitudes in Response to the Subject’s Own Name for Each Clinical Entity*


Announcement

Calendar of Events: A New Web Feature

On the new Calendar of Events site, available at http://pubs.ama-assn.org/cgi/calendarcontent and linked off the home page of the Archives of Neurology, individuals can now submit meetings to be listed. Just go to http://pubs.ama-assn.org/cgi/cal-submit/ (also linked off the Calendar of Events home page). The meetings are reviewed internally for suitability prior to posting. This feature also includes a search function that allows searching by journal as well as by date and/or location. Meetings that have already taken place are removed automatically.