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Somatosensory attention identifies both overt and covert awareness in disorders of consciousness

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Abstract

Objective: Some patients diagnosed with disorders of consciousness retain sensory and cognitive abilities beyond those apparent from their overt behaviour. Characterising these covert abilities is crucial for diagnosis, prognosis, and medical ethics. This multimodal study investigates the relationship between electroencephalographic evidence for perceptual/cognitive preservation and both overt and covert markers of awareness.

Methods: Fourteen patients with severe brain injuries were evaluated with an electroencephalographic vibrotactile attention task designed to identify a hierarchy of residual somatosensory and cognitive abilities: 1) somatosensory steady-state evoked responses, 2) bottom-up attention orienting (P3a event-related potential), and 3) top-down attention (P3b event-related potential). Each patient was also assessed with a clinical behavioural scale and two functional magnetic resonance imaging assessments of covert command following.

Results: Six patients produced only sensory responses, with no evidence of cognitive event-related potentials. A further eight patients demonstrated reliable bottom-up attention orienting responses (P3a). No patient showed evidence of top-down attention (P3b). Only those patients who followed commands, whether overtly with behaviour or covertly with functional neuroimaging, also demonstrated event-related potential evidence of attentional orienting.

Interpretation: Somatosensory attentional orienting event-related potentials differentiated patients who could follow commands from those who could not. Crucially, this differentiation was irrespective of whether command following was evident through overt external behaviour, or through covert functional neuroimaging methods. Bedside electroencephalographic methods may corroborate more expensive and challenging methods such as functional neuroimaging, and thereby assist in the accurate diagnosis of awareness.
Introduction

Disorders of consciousness (DoC) are states that a person may enter when they emerge from coma following a severe brain injury. Patients in a vegetative state (VS) do not demonstrate purposeful behaviour and are considered to lack awareness\(^1-3\). In contrast, patients in a minimally conscious state (MCS) are considered to have fluctuating awareness and demonstrate variable, but reproducible, purposeful behaviour\(^4\). Furthermore, the MCS can be sub-divided into MCS Plus or Minus on the basis of the patient’s ability to follow commands\(^5\). Patients who demonstrate accurate communication and/or functional object use are considered emergent from a MCS (EMCS)\(^4\). However, the accurate identification of a patient’s diagnostic group comprises a considerable clinical challenge\(^1-3,6-8\).

To facilitate more accurate diagnosis of the DoC, researchers have developed brain imaging paradigms to assess volition and command following in the absence of outward responsiveness\(^9-14\). Patients who produce behaviour consistent with a VS, but who exhibit evidence of covert awareness with functional neuroimaging – such as imagining movements to command\(^6,9,10,13,15-17\) – have been considered to exhibit a non-behavioural MCS\(^18\). However, in both behavioural and neuroimaging-based assessments, a patient may produce a false negative due to fatigue or insufficient cognitive resources to successfully complete the demanding diagnostic task\(^8,19\).

Researchers have developed assessments of brain function to place a patient along a hierarchy of increasingly complex attentional information processing\(^20-24\). However, there are inconsistencies in the prognostic value of the event-related potentials used in these hierarchical approaches; some investigators have reported positive prognostic value in these attentional markers\(^25\), while others have not\(^26\). These inconsistencies may have occurred because multimodal assessments were not used to identify patients in a non-behavioural MCS. Therefore, 15% of the patient sample considered to be VS may have possessed a non-behavioural MCS and consequently misrepresented the diagnostic category\(^27\). Similarly, most studies of patients with DoC employ auditory stimulation because many patients lack oculomotor control; however, this tendency limits the characterisation of a patient’s sensory abilities to the auditory domain.

We report a hierarchical cognitive assessment in a sample of fourteen patients with severe brain injuries using vibrotactile stimulation. The assessment employed an oddball paradigm to elicit steady-state evoked responses of sensory processing and event-related potential (ERP) markers of bottom-up and top-down attention (the P3a and P3b, respectively)\(^28\). As with previous hierarchical designs, this
approach discretizes a patient’s sensory and cognitive abilities. A novel aspect of our method is the assessment of a patient’s ability to sense and attend to touch. Importantly, patients were also evaluated using two previously established neuroimaging-based assessments of covert command following – mental imagery\(^6,9,10,13,15–17\) and selective auditory attention\(^29,30\) – and a clinical behavioural assessment\(^31\).

By identifying patients with covert command following abilities, these additional assessments ensured a more accurate representation of each patient’s level of awareness. Furthermore, we were in a position to test the divergence and convergence of these methods. It was expected that ERP markers of higher-order attention would be evident in patients who were aware, either expressed overtly in their behaviour, or covertly by wilful modulations of brain activity detected with neuroimaging.

**Materials and methods**

**Participants**

Fourteen patients [mean age 41 (range: 19 to 58) years] contributed sufficient data for inclusion in this investigation. Seven patients were diagnosed as VS\(^3\), four patients were diagnosed as MCS, two patients were diagnosed as EMCS\(^4\), and one patient was diagnosed with Locked-In Syndrome (LIS)\(^32\). Six patients had sustained traumatic brain injuries from motor vehicle accidents. The remaining eight patients had sustained non-traumatic brain injuries from different aetiologies including cardiac arrest (3 cases) and near-drowning (1 case; see Supplementary Table 1). Each patient’s surrogate decision maker provided informed, written consent for the patient’s participation in the study. Ethical approval was obtained from the University of Western Ontario’s Health Sciences Research Ethics Board (London, Canada).

As a scientific control, a sample of fifteen healthy volunteers also participated in the somatosensory selective attention task. These participants ranged in age from 17 to 23 years (mean age 18 years). All healthy volunteers provided informed written consent and received course credit for their participation. The Psychology Research Ethics Board of the University of Western Ontario (London, Canada) provided ethical approval for the control study. Control studies of the other neuroimaging paradigms have been reported elsewhere\(^15,30,33\).

**Procedure**

For each patient, participation in this study comprised assessments with: (1) electroencephalography (EEG) during their completion of a somatosensory selective attention paradigm;
(2) functional magnetic resonance imaging (fMRI) during their completion of a mental imagery paradigm; (3) fMRI during their completion of an auditory selective attention paradigm; and (4) the Coma Recovery Scale-Revised (CRS-R; see Supplementary Tables 1 and 2). fMRI data from Patient EMCS2 could not be analysed due to excessive motion artefacts. However, this patient was included in this investigation because his ability to follow simple commands and communicate was evident from his overt behaviour. Similarly, the data for Patient VS7 from one fMRI session (selective auditory attention) were discarded due to excessive movement. This patient was included in the current investigation because useable data were obtained from this patient for the other three paradigms.

All patients completed the two fMRI paradigms within a two-day period. Ten patients completed the fMRI assessments within two days of their EEG assessments (see Supplementary Table 2). The other four patients completed the EEG assessments after the fMRI assessment with the following delay: 1.5-months (EMCS1); 7.5-months (MCS3); 1-year (VS3); and 3.5-years (VS7). Only Patient MCS3 demonstrated a clinical status change between assessments with EEG and fMRI (MCSw to MCS+). Given the aetiology, age, and time post-ictus of those patients with a year or more between assessments (Supplementary Table 2), it is unlikely (although not impossible) that either of these patients underwent a change in their conscious states between assessments. Indeed, Patients VS3 and VS7 demonstrated overt behaviour consistent with a VS at all assessments.

**Somatosensory selective attention paradigm**

Participants completed a short somatosensory selective attention task as their EEGs were recorded. One stimulator was affixed to each wrist and the upper back (three total). Each stimulator administered non-painful vibrotactile stimuli via a motor housed in a rubberized casing. A similar paradigm has also been evaluated for patients with LIS. The experiment comprised 14 blocks. Participants were presented with a series of vibrations alternating among their wrists (10% per wrist) and upper back (80%). A vibration occurred every 200ms and lasted for 50ms. The number of vibrations presented to each wrist in a block was selected on a random uniform interval from 28 to 32. There was always a minimum of three (maximum=21) upper back stimuli between wrist vibrations; on average, 49% (standard deviation=13%) of the wrist stimuli followed exactly three upper back stimuli. Participants were instructed to count the vibrations presented only to the target wrist. The experimenter touched the patient’s target wrist after the instruction. The right wrist was always the target wrist for the first block and subsequently alternated between the left and right wrists. The healthy volunteers reported
their count at the end of each block; these participants reported the correct number of vibrations for 12/14 blocks on average (all reports were within ±3 of the true number of targets). One block of trials lasted for approximately one minute.

**Mental imagery paradigm**

During an fMRI scan, patients were asked to engage in two mental imagery paradigms. In the motor imagery task, patients were instructed to imagine swinging their right arm to hit a tennis ball. In the spatial navigation task, patients were instructed to imagine walking from room to room in their house and visualise all objects they would encounter. Instructions were delivered with noise cancellation headphones (Silent ScanTM, Avotec Inc. for patients scanned in the Trio system, as well as Patient VS6 [first visit], and Sensimetrics S14 for the patients scanned in the Prisma system, including Patient VS6 [second visit]). Patients VS1, VS2, VS4, VS5, VS6 (second visit), MCS4, and EMCS1 completed two sessions of each task, while patients VS3, VS6 (first visit), VS7, MCS1, MCS2, MCS3, and LIS1 completed only one session due to scanner availability or patient fatigue.

**Auditory selective attention paradigm**

The fMRI selective auditory attention paradigm has been previously described in healthy individuals and patients with DoC, and is designed to identify an ability to follow commands to selectively attend to stimuli – *i.e.*, top-down attention. On each trial, participants were instructed to either count a target word (‘yes’ or ‘no’) presented among pseudorandom distractors (spoken digits one to nine), or to relax. Each trial had an on/off design: sound (~22.5s) followed by silence (10s). The scan lasted five minutes, including instructions.

**Replication data**

Each task alternated five 30-second blocks of mental imagery and five 30-second blocks of rest for a total of five minutes. Patients VS4, MCS3, and EMCS1 participated in second assessments with the somatosensory selective attention task and the CRS-R. These assessments occurred from 2- to 3.5-months following their initial participation. Patient VS6 completed a second assessment with all paradigms (CRS-R, fMRI, and EEG) 22-months after her initial assessment. All four patients maintained their clinical status at follow-up (Supplementary Table 2).

**EEG data acquisition and pre-processing**
EEG data were recorded at sites FC1, Fz, FC2, C3, Cz, C4, CP1, CP2, Pz, Oz, PO7, and PO8 using an electrode cap with the g.Gamma active electrode system (g.tec Medical Engineering GmbH, Austria). This montage was selected following a previous study conducted in patients with LIS\textsuperscript{35} and previous work concerning optimal P300 classification\textsuperscript{36}. Data were sampled at 256 Hz and filtered between 0.5 and 30 Hz using a digital Butterworth filter. Stimuli were presented with the g.VIBROstim box (g.tec Medical Engineering GmbH, Austria) using a custom MATLAB\textsuperscript{®} script for Simulink\textsuperscript{®} (MathWorks, Inc., Natick, MA). The recordings were referenced to the right earlobe with a forehead (Fpz) ground. Impedances were kept below 5 kΩ. Data processing was conducted with EEGLAB\textsuperscript{37}. The data were segmented into 1-second epochs with a 200ms pre-stimulus period, and linear detrending and baseline correction were applied to each epoch. For artefact correction, all trials containing data with voltages exceeding ±100 µV were rejected. In a second step, the kurtosis of the signal across all channels was calculated for each stimulus type separately, and all trials exceeding 2.5 standard deviations of the mean were rejected. Final trial numbers are reported in (Table 1).

**fMRI data acquisition and pre-processing**

The MRI data were acquired in a 3-Tesla Siemens scanner (Siemens, Erlangen, Germany) with a Siemens 32-channel head-coil at the Centre for Functional and Metabolic Mapping at Robarts Research Institute, Western University, Canada. The patients were recruited over 30-months, in which time the 3-T scanner was upgraded. Three patients (VS3, VS7, and MCS3) were scanned in a Magnetom Trio system. All other patients were scanned in a Magnetom Prisma system. Functional echo-planar images of 36 slices covering the whole brain were acquired (repetition time=2000ms, echo time=30ms, matrix size=420 x 420, slice thickness=3 mm, in-plane resolution=3×3 mm, flip angle=78°; for patients VS6 and LIS1 only, matrix size=384x384 and flip angle=75°). High-resolution T1-weighted 3D images were acquired in the same session (Trio system: repetition time=2300ms, echo time=2.98ms, inversion time=900ms, matrix size=256×240, voxel size 1 × 1 × 1 mm, flip angle=9°; Prisma system: repetition time=2300ms, echo time=2.32ms, inversion time=900ms, matrix size=256x256, flip angle=8°; for patients VS6 and LIS1 only, matrix size=240 x 256 and flip angle=9°). Data from the mental imagery paradigm were pre-processed using SPM8 (http://www.fil.ion.ucl.ac.uk/spm), as described elsewhere\textsuperscript{13}. For the selective attention paradigm, pre-processing was performed with the AA software\textsuperscript{38}.

**Statistical analyses**
EEG responses

The EEG data were assessed for the presence of a steady-state evoked potential to the repetitive vibrotactile stimulation. As one vibration occurred every 200ms, an evoked response was considered present when the averaged peak of the frequency spectrum of the data at the stimulation rate (5 Hz) and its first harmonic (10 Hz) was significantly higher than the background noise. A frequency spectrum was calculated with a discrete Fourier transform over the entire 1-second epoch from the average of all trials using data only from site Pz. An $F$ ratio ($\alpha=.05; F_{2,20}=3.49$) was computed to compare the power at 5 and 10 Hz with the average power in the ten adjacent ~1 Hz frequency bins (2-4 Hz, 6-9 Hz, and 11-13 Hz).

Two analyses of the EEG data were conducted to identify the attention-based event-related potentials. For the bottom-up attention effect (P3a), responses to wrist (deviant) and upper back (standard) stimuli were compared. A random subset of the standard stimuli (equal in number to the deviant stimuli) was selected because there were many more standard than deviant stimuli. For the top-down attention effect (P3b), responses to the target and non-target wrist stimuli were compared. Trial numbers were matched between the target and non-target trials. Data from 50 to 750ms post-stimulus were analysed using the cluster-mass procedure of the MATLAB® toolbox FieldTrip. This technique has been described in detail previously. In the first step, data were compared at each time-point using a $t$-test. In the second step, $t$-values of adjacent spatiotemporal points with $p<.05$ were clustered together by summatng their $t$-values. The largest cluster was retained. This entire procedure was repeated 1000 times with recombination and randomized resampling of the ERP data. This Monte Carlo method generated a nonparametric estimate of the $p$-value representing the statistical significance of the originally identified cluster.

Blood oxygen level-dependent (BOLD) mental imagery responses

Single subject fixed-effect analyses were performed for each patient. The analysis was based on the general linear model using the canonical hemodynamic response function implemented with SPM8 (http://www.fil.ion.ucl.ac.uk/spm). The analysis pipeline was previously reported. Linear contrasts were used to obtain subject-specific estimates, and results were thresholded at a voxel level, familywise error (FWE), whole-brain $p<.05$. When no significant activations were found at this level, the statistical threshold was reduced to an uncorrected $p<.001$ because of the strong anatomical a priori
hypotheses. This less conservative threshold excluded the possibility of failing to detect more subtle changes in the signal.

**BOLD auditory selective attention responses**

The general linear model (SPM8) was used to explore effects of interest. Two event types were defined corresponding to the on/off periods (count/relax; ~22.5s, or vice-versa). The silent period (10s) served as an implicit baseline for all trials. Events for these regressors were modelled by convolving boxcar functions with the canonical hemodynamic response function. Also included in the general linear model were the following nuisance variables: the movement parameters in the three directions of motion and three degrees of rotation, and the mean of each scan. Linear contrasts were used to obtain subject-specific estimates for the effect of interest. Clusters that survived the $p<.05$ threshold after the FWE correction were reported as significant.

**Results**

All patient outcomes are summarized in (Figure 1) and (Supplementary Table 3).

**EEG responses**

A steady-state evoked potential was detected in the EEG data of all patients ($n=14$) and all healthy volunteers ($n=15$; Figure 2).

**Bottom-up attention effects** (deviant versus standard stimuli) were detected from eight patients and all of the healthy volunteers ($n=15$; Figure 3). All patients who demonstrated a differential response to the deviant versus standard stimuli also demonstrated evidence of command following in either a behavioural or a neuroimaging-based assessment (Figure 1 and Supplementary Table 3).

Top-down ERP attention effects (target versus non-target wrist vibrations) were not detected from any of the patients. However, this ERP effect was evident for healthy volunteers at the group level ($n=15$) and at the single-subject level, albeit with a hit-rate of 67% (Figure 4). Hit-rates of at least 80% (12/15) and 100% (15/15) have been reported for fMRI-detected mental imagery and selective attention respectively. Given the relatively lower sensitivity of the top-down attention ERP analysis (*i.e.*, 67%), additional post-hoc comparisons were conducted. While the number of trials available after artefact rejection did not differ across groups (Table 1; $\chi^2(2)=0.21, p=0.9$), some patients had many fewer trials available than healthy individuals. The single-subject ERP analyses for the healthy volunteers were thus
repeated in the post-hoc analyses using only a pseudorandom subset of trials equal in number to the minimum number of trials available in the single-subject analyses of the patient data (180 trials, in the case of Patient MCS2).

Bottom-up attentional ERP effects were detected at the single-subject level for all healthy volunteers when as few as 180 trials were included for each stimulus type. However, top-down attentional ERP effects were detected from only seven healthy volunteers. Subsequent analyses revealed that a minimum of 300 trials were required to detect the top-down attentional ERP effects from the same 10 healthy volunteers as in the a priori analyses. Four patients did not have enough trials available to meet this criterion. Overall, these analyses indicate that the top-down attentional ERP effect may not have been detected in some single-subject analyses due to low trial numbers. Nevertheless, the bottom-up attentional ERP effect was robust to data loss.

BOLD mental imagery responses

In her first visit, Patient VS6 produced reliable, appropriate activation during the motor imagery task in the supplementary motor area and cerebellum bilaterally at an uncorrected $p<.001$ (cluster level FWE-corrected $p<.05$). In her second visit, Patient VS6 produced reliable, isolated clusters of activation during the motor imagery and spatial navigation tasks in the left precentral gyrus at an uncorrected $p<.001$ (cluster level FWE-corrected $p<.05$). The patient was thus reclassified as in a non-behavioural MCS$^{18}$.

Patients VS7 showed high levels of motion requiring 37% and 37.5% of his data to be discarded (for motor imagery and spatial navigation respectively). The analysis of the remaining data revealed appropriate activation during the spatial navigation task only (i.e., the left occipito-parietal junction at uncorrected $p<.001$). The patient was thus reclassified as in a non-behavioural MCS$^{18}$.

Patients MCS3, MCS4, EMCS1, and LIS1 showed reliable activation during the spatial navigation task only. This involved: bilateral occipito-parietal junction (uncorrected $p<.001$) for MCS3; right temporo-occipito-parietal junction (FWE-corrected $p<.05$), as well as right dorsal premotor cortex, right insular cortex, and right putamen (uncorrected $p<.001$) for MCS4; right occipito-parietal junction, a region in the boundaries between right lingual gyrus/parahippocampal cortex, left precentral gyrus (comprising the supplementary and pre-supplementary motor areas), as well as some less typical areas such as the inferior frontal gyrus, the left superior temporal gyrus, and the left striatum (FWE-corrected
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$p<.05$) for EMCS1; and supplementary motor area, right precentral gyrus, occipito-parietal junction, posterior temporoo-occipital region, and the cerebellum (uncorrected $p<.001$) for LIS1.

The remaining seven patients (VS1-5, MCS1, and MCS2) showed no activation at the conservative FWE-corrected statistical threshold, or at uncorrected $p<.001$.

**BOLD auditory selective attention responses**

Of the patients diagnosed as in a VS, only Patient VS6 showed significantly more activation following the instruction to count than to relax. This patient showed significant activation in the temporal and parietal cortex bilaterally (FWE-corrected at $p<.05$).

Patients MCS1-4 and LIS1 also showed significantly more activation following the instruction to count than to relax. Patient MCS1 showed significant activation in the frontotemporal and parietal cortex bilaterally. Patient MCS2 showed significant activation in the temporal cortex bilaterally (FWE-corrected at $p<.05$). Patient MCS3 showed significant activation in the parietal cortex bilaterally. Patient MCS4 showed significant activation in the frontotemporal and parietal cortex bilaterally (FWE-corrected at $p<.05$). Patient LIS1 produced significant brain activity in the frontotemporal cortex bilaterally (FWE-corrected at $p<.05$).

Of note, Patient EMCS1 did not show significant differences in activation in the command following task even though she was able to follow commands with her overt behaviour immediately prior to her assessment. Patients VS7 and EMCS2 were excluded from this analysis because both patients moved excessively during their functional scans.

**Correspondence between command following and EEG responses**

The main hypothesis in this investigation was that patients who were aware would exhibit concordant EEG markers of higher-order attention processing. While top-down processing (P3b) was not detected from any patients, an interesting observation from the current data is the relationship between a specific marker of awareness – command-following – and the bottom-up attention orienting ERP effect, the P3a. A patient was considered to have evidence of such awareness if they demonstrated evidence of command following in any one of the three non-EEG assessments (selective auditory attention, mental imagery, or a behavioural assessment with the CRS-R). This approach is consistent with clinical behavioural guidelines in which a diagnosis of awareness (MCS) is given if a patient follows commands on one occasion across multiple assessments. A Fisher’s exact test revealed a
significant positive association between evidence for command following and evidence for the P3a (\(p = .007\); note \(p = .0047\) if the two observations of Patient VS6 are not included to maintain the assumption of independence). This relationship is summarised in (Figure 1).

Replication data

The replication results are depicted in (Figure 5). All patients exhibited consistent effects across assessments with the exception of Patient VS6 for whom a P3a was significant only during her initial assessment.

Discussion

We investigated a novel EEG method for the assessment of residual sensory and cognitive processing alongside two fMRI-based assessments of covert command following and one behavioural assessment of overt command following in a sample of fourteen patients with severe brain injuries. The primary novel finding of this work is the relationship between an ERP marker of bottom-up attention orienting (the P3a) and command following such that all patients with a P3a response demonstrated positive evidence of command following. Similarly, most patients who did not generate a P3a response also did not demonstrate evidence of command following (see Figure 1 and Supplementary Table 3).

Some investigators have reported positive prognostic value in the presence of a P300 following traumatic brain injury\(^25\). There have also been reports of correlations between cognitive ERPs and behavioural markers of awareness\(^14,24\), as well as the prediction of recovery from the DoC using cognitive ERPs\(^47,26\). Crucially, the current study included two neuroimaging-based assessments of covert command following. This step is important given that a recent meta-analysis estimates a 15% rate of covert awareness among patients diagnosed as in a VS\(^27\). Previous studies of the P300 in patients with DOC are likely to have included patients capable of covert command following, thus obscuring the relationship reported here. While the feasibility of routine neuroimaging assessments in clinical practice is limited by important health, safety, and financial factors, the findings of this work suggest that these assessments are necessary to elucidate the relationship between a patient’s conscious state and their residual sensory and cognitive abilities.

It is curious that an ERP marker of unconscious (or preconscious) processing – i.e., the P3a – is closely linked to awareness in this work. Indeed, the P3a can be elicited by unattended stimuli and during REM sleep and deep sedation\(^28,48\). We speculate that the correspondence between the P3a and
command following stems from the overlap of the neural networks that support attention, and those that are relatively more preserved in conscious patients\(^{49,50}\). Indeed, frontal lobe lesions have been associated with diminished P3a responses to auditory\(^{51}\) and somatosensory\(^{52}\) stimulation. Equally, this association suggests that a P3a response may be less informative for patients with specific frontal lobe injuries. Nevertheless, a P3a can be elicited without the explicit collaboration of the individual – \textit{i.e.}, without following task instructions\(^{48}\). This feature is appealing, as it suggests that a passive assessment of attention orienting, which entails lower cognitive demands than active assessments of voluntary top-down attention, may be sufficient to identify patients with covert awareness.

The P3b marker of top-down attention in the current EEG task was not detected from any of the patients in this sample, as has been reported previously\(^{53}\). In fact, P3b responses in the current work were detected from only 67\% (10/15) of the healthy volunteers. \textit{Post-hoc} analyses of the ERP data indicated that this low sensitivity may be exacerbated by the fewer usable trials in the patient data, as this comparison was sensitive to a reduced signal-to-noise ratio. Additionally, time-variant levels of arousal and fatigue characteristic of the DoC may have led to inconsistent engagement in the counting task needed to generate the top-down ERP effect\(^{8,19}\). In contrast to the fMRI-based selective attention task, the selective attention manipulation in the EEG task may have placed higher cognitive demands on participants due to the longer duration of the EEG task. Participants were required to sustain attention for five minutes in \textasciitilde 22.5-second blocks for both fMRI tasks, whereas the EEG task involved fifteen minutes of attention in \textasciitilde 1-minute blocks. The EEG task was longer to ensure that a high EEG signal-to-noise ratio was achieved, and \textit{post-hoc} analyses confirmed that the top-down ERP effect was sensitive to trial numbers. Unfortunately, increased task duration requires participants to sustain attention for an even longer period, making it unlikely that this manipulation would increase the sensitivity of the task.

Some investigators use machine learning to circumvent these issues and address possible spatiotemporal variations in the electrocortical responses of patients with brain injuries\(^{54}\). For simplicity of interpretation and consistency with clinical methods, we employed a more traditional approach to comparing scalp voltages. While no false alarms were evident in the current sample, misses occurred with two patients – \textit{i.e.}, patients demonstrated evidence of command following but no evidence of a P3a. As has been discussed elsewhere, signs of awareness in both behavioural and neuroimaging assessments may be missed due to fluctuating arousal\(^{13}\). Nevertheless, when a P3a is elicited, the current data suggest the sophisticated cognitive networks that underlie an ability to follow commands are also preserved.
The detection of awareness in the DoC is a clinical standard of care. In order to provide sufficient evidence to influence clinical practice, it is essential to compare novel assessments to existing techniques. The current investigation allowed for a comparison of two previously reported neuroimaging-based assessments of covert command following, based on mental imagery and selective auditory attention. The results of these assessments converged for nine of the twelve patients with useable data from both paradigms. Two patients demonstrated positive evidence of command following in only the selective auditory attention task, while one patient showed positive evidence of command following only in the mental imagery task. The behavioural profile of the DoC – that is, time-variant fatigue and arousal – always affords the possibility that a patient did not demonstrate positive evidence of covert command following due to lack of voluntary engagement in the task. Likewise, false negatives occur in assessments of healthy volunteers. Nevertheless, the less than perfect correspondence of the two covert fMRI command following tasks may have occurred because the demands of one task were better suited to the patient. For example, some individuals find it difficult to engage in motor imagery, and in some reports, brain-computer interfaces based on selective attention tasks are successfully operated by more users than those based on responses to motor imagery. Accordingly, assessments of covert command following based on selective attention may be better suited to a general population. Overall, however, an optimal evaluation of a patient with a DoC should include multiple assessments to maximise the likelihood of detecting responses that are not evident from overt behaviour. In the absence of unambiguous ground truth, an investigation of the concordance between assessments may be the best way to improve diagnostic and prognostic accuracy.

In summary, the brain responses of fourteen patients with severe brain injuries were assessed using an EEG-based somatosensory selective attention task, two fMRI-based assessments of covert command following, and one behavioural instrument. While limited by a relatively small sample of patients, the data tentatively suggest that the detection of a somatosensory bottom-up P3a effect in a patient correlates with an ability to follow commands, as evaluated by multimodal assessments. This provides evidence that a bedside somatosensory oddball procedure can improve diagnostic accuracy in the DoC and more accurately characterise the level of neurocognitive preservation. Overall, this work provides a valuable addition to neuroimaging batteries for the clinical assessment of patients with DoC and convergent, multimodal evidence for the utility of these techniques.
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Author Contributions

R.M.G., D.F.E., L.N., and D.C. contributed to conception and design of the study. R.M.G., D.F.E., L.N., and D.C. contributed to data collection and analysis. R.M.G., D.C., S.C., and A.M.O contributed to writing the manuscript.

Potential Conflicts of Interest

Nothing to report.
References


Figure Captions

Figure 1. Summary of the relationship between command following and outcomes on the selective somatosensory attention task.

The summary depicts the number of patients and healthy volunteers who generated each of the three possible outcomes on the somatosensory selective attention task.

VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

Figure 2. Steady-state evoked responses to the repetitive vibrotactile stimulation.

Power spectra (top panels) and averaged EEG responses (bottom panels) calculated over a period of 1-second. Analyses were conducted using the data recorded from site Pz only; each waveform (bottom panels) is depicted with ±1 standard error of the mean.

EEG=electroencephalography; **=p<0.01; ***=p<.001; VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

Figure 3. Bottom-up attention event-related potentials to the standard and deviant vibrotactile stimulation.

Spatiotemporal clusters were calculated across all twelve electrodes and are depicted with ±1 standard error of the mean in colour-matched shading. The electrodes included in the significant spatiotemporal cluster are enclosed with a black line on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. (A) depicts the grand-averaged ERP effect for the healthy volunteers, (B) depicts the single-subject ERP effects for the healthy volunteers (p<9.9E-03 in all cases), and (C) depicts the single-subject ERP effects for the patients with statistically significant results.

VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

Figure 4. Top-down attention event-related potentials to the target and non-target vibrotactile stimulation for the healthy volunteers.
Spatiotemporal clusters were calculated across all twelve electrodes with each waveform depicted with ±1 standard error of the mean. The electrodes included in the significant spatiotemporal cluster are enclosed with a black outline on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. The grand-averaged result ($n=15$) is depicted in (A). For the single subject results (B), only results from participants with statistically significant clusters are shown.

Figure 5. Replication data from the four patients with whom follow-up investigations were conducted.

Data are depicted for the initial and follow up tests of Patients VS4, MCS3, EMCS1, and VS6, as labelled. For the steady-state evoked potentials, power spectra (top left panels within each cell) and averaged EEG data (bottom left panels within each cell) were calculated over a period of 1-second. Analyses were conducted using the data recorded from site Pz only; each waveform is depicted with ±1 standard error of the mean. For the bottom-up attention ERP effects (right panels within each cell), spatiotemporal clusters were calculated across all twelve electrodes and are depicted with ±1 standard error of the mean. The electrodes included in the significant spatiotemporal cluster are enclosed with a black line on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. For Patient VS6 only, two separate fMRI assessments were conducted at each testing session. For the fMRI mental imagery paradigm, significant task-related fMRI activation is depicted (Imagery>Rest), and results are thresholded at an uncorrected $p<.001$. For the fMRI selective auditory attention task, only activation clusters within the attention network (Count>Relax) that survived the familywise error correction threshold of $p<.05$ at the whole-brain level are displayed. The fMRI results are rendered on the patient’s T1 anatomical MRI image, and scales depicting the $t$-value statistical maps are inset.

* = $p<0.05$; ** = $p<0.01$; *** = $p<.001$; n.s. = not statistically significant; VS = vegetative state; MCS = minimally conscious state; EMCS = emergent from a minimally conscious state.
Table 1. Number of trials available for the analyses of the EEG data from the somatosensory selective attention paradigm following artefact rejection.

<table>
<thead>
<tr>
<th>Stimulus Type&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Upper Back</th>
<th>Target Wrist</th>
<th>Non-Target Wrist</th>
<th>Trials Rejected (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M (MIN-MAX)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients (n=14)</td>
<td>2614 (1591-3246)</td>
<td>313 (188-384)</td>
<td>311 (180-388)</td>
<td>35 (20-59)</td>
</tr>
<tr>
<td>Controls (n=15)</td>
<td>2890 (2718-5026)</td>
<td>345 (327-363)</td>
<td>345 (321-359)</td>
<td>25 (20-32)</td>
</tr>
</tbody>
</table>

<sup>Note</sup>. M=mean; MIN=minimum; MAX=maximum.

<sup>a</sup>A 2x3 Chi-square goodness of fit test indicated that the minimum number of trials in each of the three stimulus types did not significantly differ between the controls and patients, $\chi^2(2)=0.21$, $p=0.9$. 
Summary of the relationship between command following and outcomes on the selective somatosensory attention task.

The summary depicts the number of patients and healthy volunteers who generated each of the three possible outcomes on the somatosensory selective attention task.

VS = vegetative state; MCS = minimally conscious state; EMCS = emergent from a minimally conscious state; LIS = Locked-In Syndrome.

80x60mm (300 x 300 DPI)
Command-following

- Overt (behaviour)
- Covert (fMRI)

Healthy 5 10
LIS 1 2
EMCS 1 3
MCS 5 2
VS

Sensory Bottom-up Top-down attention

Proportion within each row

(Black and white version of Figure 1 for print)
80x60mm (300 x 300 DPI)
Figure 2. Steady-state evoked responses to the repetitive vibrotactile stimulation. Power spectra (top panels) and averaged EEG responses (bottom panels) calculated over a period of 1-second. Analyses were conducted using the data recorded from site Pz only; each waveform (bottom panels) is depicted with ±1 standard error of the mean.

EEG=electroencephalography; **=p<0.01; ***=p<.001; VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

170x427mm (300 x 300 DPI)
(A) Steady-state evoked potentials for the patients.

VS1 VS2 VS3 VS4 VS5

VS6 VS7 MCS1 MCS2

MCS3 MCS4 EMS1 EMS2 LIS1

(B) Steady-state evoked potentials for the healthy volunteers.

Control 1 Control 2 Control 3 Control 4 Control 5

Control 6 Control 7 Control 8 Control 9 Control 10

Control 11 Control 12 Control 13 Control 14 Control 15

(Black and white version of Figure 2 for print)

170x427mm (300 x 300 DPI)
Figure 3. Bottom-up attention event-related potentials to the standard and deviant vibrotactile stimulation. Spatiotemporal clusters were calculated across all twelve electrodes and are depicted with ±1 standard error of the mean in colour-matched shading. The electrodes included in the significant spatiotemporal cluster are enclosed with a black line on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. (A) depicts the grand-averaged ERP effect for the healthy volunteers, (B) depicts the single-subject ERP effects for the healthy volunteers (p<9.9E-03 in all cases), and (C) depicts the single-subject ERP effects for the patients with statistically significant results. VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

170x160mm (300 x 300 DPI)
(A) Grand-averaged bottom-up attention ERP effect for the healthy volunteers (n=15).

(B) Bottom-up attention ERP effects for the healthy volunteers.

(C) Bottom-up attention ERP effects for the patients with statistically significant results.

(Black and white version of Figure 3 for print)
170x160mm (300 x 300 DPI)
Figure 4. Top-down attention event-related potentials to the target and non-target vibrotactile stimulation for the healthy volunteers. Spatiotemporal clusters were calculated across all twelve electrodes with each waveform depicted with ±1 standard error of the mean. The electrodes included in the significant spatiotemporal cluster are enclosed with a black outline on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. The grand-averaged result (n=15) is depicted in (A). For the single subject results (B), only results from participants with statistically significant clusters are shown.

170x262mm (300 x 300 DPI)
(A) Grand-averaged top-down attention ERP effect for the healthy volunteers (n=15).

(B) Top-down attention ERP effects for the healthy volunteers with statistically significant results.

(Black and white version of Figure 4 for print)
170x262mm (300 x 300 DPI)
Figure 5. Replication data from the four patients with whom follow-up investigations were conducted. Data are depicted for the initial and follow-up tests of Patients VS4, MCS3, EMCS1, and VS6, as labelled. For the steady-state evoked potentials, power spectra (top left panels within each cell) and averaged EEG data (bottom left panels within each cell) were calculated over a period of 1-second. Analyses were conducted using the data recorded from site Pz only; each waveform is depicted with ±1 standard error of the mean.

For the bottom-up attention ERP effects (right panels within each cell), spatiotemporal clusters were calculated across all twelve electrodes and are depicted with ±1 standard error of the mean. The electrodes included in the significant spatiotemporal cluster are enclosed with a black line on each topographic plot. The temporal boundaries and the probability value of each cluster are indicated with shading and inset text. For Patient VS6 only, two separate fMRI assessments were conducted at each testing session. For the fMRI mental imagery paradigm, significant task-related fMRI activation is depicted (Imagery>Rest), and results are thresholded at an uncorrected p<.001. For the fMRI selective auditory attention task, only activation clusters within the attention network (Count>Relax) that survived the familywise error correction threshold of p<.05 at the whole-brain level are displayed. The fMRI results are rendered on the patient’s T1 anatomical MRI image, and scales depicting the t-value statistical maps are inset.

* = p<0.05; ** = p<0.01; *** = p<.001; n.s. = not statistically significant; VS = vegetative state;
MCS=minimally conscious state; EMCS=emergent from a minimally conscious state.
170x191mm (300 x 300 DPI)
For Peer Review (Black and white version of Figure 5 for print)

VS4

MCS3

EMCS1

VS6

(Black and white version of Figure 5 for print)
170x191mm (300 x 300 DPI)
**Supplementary Table 1.** Summary of the patients recruited for this investigation, including their age, *post ictus* interval, and behaviour as measured by the Coma Recovery Scale-Revised.

All behavioural data reported here correspond with the patient’s abilities at the time when they generated the highest total score on the Coma Recovery Scale-Revised; these assessments occurred prior to the patient’s participation in the current investigation in some cases.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Diagnosis</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Aetiology</th>
<th>Post ictus (years)</th>
<th>Sensory Responses</th>
<th>Attention Responses</th>
<th>Command Following Responses</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS1</td>
<td>Non-traumatic brain injury secondary to Commodio Cordis</td>
<td>19</td>
<td>M</td>
<td>Non-traumatic brain injury secondary to Commodio Cordis</td>
<td>4.0</td>
<td>Auditory startle, Localization to sound, Visual startle, Abnormal posturing, Flexion withdrawal, Oral reflexive movement</td>
<td>Present Eye opening without stimulation Absent Absent</td>
<td>Absent Absent Absent</td>
</tr>
<tr>
<td>VS2</td>
<td>Massive myocardial infarction</td>
<td>51</td>
<td>F</td>
<td>Massive myocardial infarction</td>
<td>0.9</td>
<td>Auditory startle, Abnormal posturing, Oral reflexive movement</td>
<td>Present Eye opening with stimulation Absent Absent</td>
<td>Absent Absent Absent</td>
</tr>
<tr>
<td>VS3</td>
<td>Non-traumatic brain injury secondary to cardiac arrest</td>
<td>57 (fMRI) 58 (EEG)</td>
<td>M</td>
<td>Non-traumatic brain injury secondary to cardiac arrest</td>
<td>3.1 (fMRI) 4.1 (EEG)</td>
<td>Auditory startle, Abnormal posturing, Flexion withdrawal, Oral reflexive movement</td>
<td>Present Eye opening without stimulation Absent Absent</td>
<td>Absent Absent Absent</td>
</tr>
<tr>
<td>VS4</td>
<td>Non-traumatic brain injury secondary to</td>
<td>42</td>
<td>F</td>
<td>Non-traumatic brain injury secondary to</td>
<td>4.3</td>
<td>Auditory startle, Localization to</td>
<td>Present Eye opening without stimulation Absent Absent</td>
<td>Absent Absent Absent</td>
</tr>
<tr>
<td>VS5</td>
<td>VS</td>
<td>52</td>
<td>F</td>
<td>Hypoxic ischemic encephalopathy, severe generalized atrophy/cardiac arrest</td>
<td>Auditory startle Abnormal posturing Flexion withdrawal Oral reflexive movement</td>
<td>Present</td>
<td>Eye opening with stimulation Absent Absent Absent</td>
<td>Absent Absent Absent</td>
</tr>
<tr>
<td>VS6</td>
<td>VS/MCS*</td>
<td>44 (Test 1) 46 (Test 2)</td>
<td>F</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>Auditory startle Abnormal posturing Flexion withdrawal Oral reflexive movement</td>
<td>Present</td>
<td>Eye opening without stimulation Absent Present (Test 1) Absent (Test 2)</td>
<td>Absent Present Present</td>
</tr>
<tr>
<td>VS7</td>
<td>VS/MCS*</td>
<td>23 (MRI) 26 (EEG)</td>
<td>M</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>Auditory startle Visual startle Abnormal posturing Flexion withdrawal Oral reflexive movement</td>
<td>Present</td>
<td>Eye opening without stimulation Absent Present</td>
<td>Absent Present Present</td>
</tr>
<tr>
<td>MCS1*</td>
<td>MCS-</td>
<td>40</td>
<td>M</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>Auditory startle Visual startle Abnormal posturing Oral reflexive movement</td>
<td>Present</td>
<td>Eye opening with stimulation Visual fixation and pursuit Absent</td>
<td>Absent Absent Present</td>
</tr>
<tr>
<td>MCS2</td>
<td>MCS+</td>
<td>35</td>
<td>M</td>
<td>Non-traumatic brain injury secondary to cardiac arrest</td>
<td>Auditory startle Localization to sound Visual startle and fixation* Abnormal posturing</td>
<td>Present</td>
<td>Eye opening without stimulation Visual pursuit Present</td>
<td>Object localisation Reproducible movement to command Absent Present</td>
</tr>
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</tr>
<tr>
<td>MCS3</td>
<td>47</td>
<td>F</td>
<td>Present</td>
<td>Non-traumatic brain injury from near-drowning</td>
<td>Flexion withdrawal, Oral reflexive movement, Vocalization/Oral movement</td>
<td>Present</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>MCS4</td>
<td>25</td>
<td>F</td>
<td>Absent</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>Auditory startle, Localization to sound, Visual startle, Abnormal posturing, Flexion withdrawal, Oral reflexive movement</td>
<td>Present</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>EMCS1</td>
<td>49</td>
<td>F</td>
<td>Absent</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>Auditory startle, Localization to sound, Visual startle, Abnormal posturing, Flexion withdrawal, Oral reflexive movement, Vocalization/Oral movement</td>
<td>Present</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>Patient</td>
<td>Status</td>
<td>Age</td>
<td>Sex</td>
<td>Condition</td>
<td>CRS-R</td>
<td>EEG</td>
<td>fMRI</td>
<td>Notes</td>
</tr>
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</tr>
<tr>
<td>EMCS2</td>
<td>EMCS</td>
<td>32</td>
<td>M</td>
<td>Traumatic brain injury secondary to motor vehicle accident</td>
<td>4.1</td>
<td></td>
<td></td>
<td>(MRI data could not be acquired because the patient moved excessively during the scan)</td>
</tr>
<tr>
<td>LIS1</td>
<td>LIS</td>
<td>55</td>
<td>M</td>
<td>Brainstem infarct related to vertebral artery thrombosis</td>
<td>1.5</td>
<td></td>
<td></td>
<td>Sustained visual fixation does not necessarily reflect higher order cortical brain function in patients with DoC and non-traumatic aetiology.</td>
</tr>
</tbody>
</table>

**Notes.**

*Patient MCS1 scored in the VS range immediately prior to his participation in the EEG assessment. However, he scored in the MCS- range in another assessment with the Coma Recovery Scale-Revised several hours prior to his participation in the EEG assessment and has thus been classified as in a MCS-.*

*Sustained visual fixation does not necessarily reflect higher order cortical brain function in patients with DoC and non-traumatic aetiology.*
**Supplementary Table 2.** Coma Recovery Scale-Revised scores for each patient immediately prior to their assessment with electroencephalography and functional magnetic resonance imaging for this investigation.

<table>
<thead>
<tr>
<th>Patient</th>
<th>CRS-R Sub-scores at EEG assessment</th>
<th>CRS-R Sub-scores at fMRI assessment</th>
<th>Time between EEG and fMRI assessments</th>
<th>Time between fMRI assessments</th>
<th>Time between EEG replication sessions</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS1</td>
<td>1-Auditory startle 0-None 2-Flexion withdrawal 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 5</td>
<td>1-Auditory startle 1-Visual startle 0-None 1-Oral reflexive movement 0-None 2-Eye opening without stimulation 5</td>
<td>2 days</td>
<td>2 days</td>
<td>N/A</td>
</tr>
<tr>
<td>VS2</td>
<td>1-Auditory startle 0-None Abnormal posturing 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 4</td>
<td>1-Auditory startle 0-None 1-Abnormal posturing 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 4</td>
<td>1 day</td>
<td>2 days</td>
<td>N/A</td>
</tr>
<tr>
<td>VS3</td>
<td>1-Auditory startle 0-None 2-Flexion withdrawal 1-Oral reflexive movement 0-None 2-Eye opening without stimulation 6</td>
<td>1-Auditory startle 1-Visual startle 2-Flexion withdrawal 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 6</td>
<td>&lt;1 hour</td>
<td>1 day</td>
<td>1 year</td>
</tr>
<tr>
<td>VS4</td>
<td>1-Initial Localization to sound 1-Visual startle 2-Flexion withdrawal 1-Oral reflexive movement 0-None 2-Eye opening without stimulation 8</td>
<td>1-Auditory startle 1-Visual startle 2-Flexion withdrawal 1-Oral reflexive movement 0-None 2-Eye opening without stimulation 7</td>
<td>N/A</td>
<td>1 day</td>
<td>3.5 months</td>
</tr>
<tr>
<td>VS5</td>
<td>1-Auditory startle 0-None Abnormal posturing 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 4</td>
<td>1-Auditory startle 0-None 2-Flexion withdrawal 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 5</td>
<td>1 day</td>
<td>1 day</td>
<td>N/A</td>
</tr>
<tr>
<td>VS6</td>
<td>1-Auditory startle 0-None 2-Flexion withdrawal 1-Oral reflexive movement 0-None 1-Eye opening with stimulation 5</td>
<td>0-None 1-Visual startle 0-None 0-None 0-None 2-Eye opening without stimulation 3</td>
<td>&lt;1 hour</td>
<td>1 day</td>
<td>22 months</td>
</tr>
</tbody>
</table>

*Note: fMRI assessments conducted only at EEG Session 2.*
<table>
<thead>
<tr>
<th>VS7</th>
<th>2-Eye opening without stimulation</th>
<th>1-Auditory startle</th>
<th>0-None</th>
<th>2-Flexion withdrawal</th>
<th>1-Oral reflexive movement</th>
<th>0-None</th>
<th>2-Eye opening without stimulation</th>
<th>0-None</th>
<th>0-None</th>
<th>0-None</th>
<th>0-None</th>
<th>0-None</th>
<th>0-None</th>
<th>2-Eye opening without stimulation</th>
<th>&lt;1 hour (Same scan)</th>
<th>2 days</th>
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<tr>
<td></td>
<td></td>
<td>1-Auditory startle</td>
<td>0-None</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>&lt;1 hour (Same scan)</td>
<td>2 days</td>
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</tr>
<tr>
<td>MCS1*</td>
<td>1-Auditory startle</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>1-Eye opening with stimulation</td>
<td>5</td>
<td>1-Auditory startle</td>
<td>3-Visual pursuit</td>
<td>1-Abnormal posturing</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>1-Eye opening with stimulation</td>
<td>&lt;1 hour (Same scan)</td>
<td>2 days</td>
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<tr>
<td>MC2</td>
<td>1-Auditory startle</td>
<td>1-Visual pursuit</td>
<td>3-Visual opening</td>
<td>0-None</td>
<td>1-Eye opening with stimulation</td>
<td>8</td>
<td>1-Auditory startle</td>
<td>3-Visual pursuit</td>
<td>4-Object manipulation</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>1-Eye opening with stimulation</td>
<td>&lt;1 hour (Same scan)</td>
<td>2 days</td>
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<tr>
<td>EM2</td>
<td>1-Auditory startle</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>8</td>
<td>1-Auditory startle</td>
<td>3-Visual pursuit</td>
<td>1-Abnormal posturing</td>
<td>1-Oral reflexive movement</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>&lt;1 hour (Same scan)</td>
<td>2 days</td>
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<tr>
<td>EMCS1</td>
<td>1-Auditory startle</td>
<td>4-Object recognition</td>
<td>6-Functional object use</td>
<td>2-Vocalization/Oral movement</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>0-None</td>
<td>2-Eye opening without stimulation</td>
<td>&lt;1 hour (Same scan)</td>
<td>2 days</td>
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<tr>
<td>EMCS2</td>
<td>1-Auditory startle</td>
<td>4-Object recognition</td>
<td>6-Functional object use</td>
<td>3-Intelligible Verbalization</td>
<td>2-Functional: Accurate</td>
<td>3-Attention</td>
<td>4-Object recognition</td>
<td>5-Object recognition</td>
<td>5-Automatic motor response</td>
<td>0-None</td>
<td>2-Functional: Accurate</td>
<td>3-Attention</td>
<td>1 day</td>
<td>2 days</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(Imagery) (Selective Attention) (Replication) (MRI assessments conducted once 7.5 months prior to EEG Session 1)
<table>
<thead>
<tr>
<th></th>
<th></th>
<th>2 (Selective Attention)</th>
<th>4- Consistent movement to command</th>
<th>5- Object recognition to command</th>
<th>5- Automatic motor response</th>
<th>3- Intelligible Verbalization</th>
<th>2- Functional: Accurate</th>
<th>3- Attention</th>
<th>22</th>
<th>1 day</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>LIS1</td>
<td>1</td>
<td>4- Consistent movement to command</td>
<td>5- Object recognition</td>
<td>0- None</td>
<td>1- Oral reflexive movement</td>
<td>2- Functional: Accurate</td>
<td>3- Attention</td>
<td>15</td>
<td>1 (Both paradigms)</td>
<td>4- Consistent movement to command</td>
<td>5- Object recognition</td>
</tr>
</tbody>
</table>

CRS-R=Coma Recovery Scale-Revised; EEG=electroencephalography; fMRI=functional magnetic resonance imaging; VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome.

Notes: 1Patient MCS1 scored in the VS range immediately prior to his participation in the EEG assessment. However, he scored in the MCS- range in another assessment with the Coma Recovery Scale-Revised several hours prior to his participation in the EEG assessment and has thus been classified as in a MCS-.
2Patient EMCS2 was not assessed with the CRS-R at her replication session. However, she was able to communicate using an arm movement.
**Supplementary Table 3.** Patient outcomes on the behavioural, neuroimaging-based, and electroencephalography-based assessments.

Only positive results are depicted for the EEG- and fMRI-based assessments. For the fMRI mental imagery paradigm, significant task-related fMRI activation is labeled by region (Imagery>Rest), and results are thresholded at an uncorrected $p<.001$. For the fMRI selective auditory attention task, only activation clusters within the attention network (Count>Relax) that survived the familywise error correction threshold of $p<.05$ at the whole-brain level are displayed. All fMRI results are rendered on each patient’s T1 anatomical MRI image, and scales depicting the $t$-value statistical maps are inset.

EEG=electroencephalography; fMRI=functional magnetic resonance imaging; SMA=supplementary motor area; OPJ=occipito-parietal junction; TOPJ=temporo-occipito-parietal junction; PHC=parahippocampal cortex; IFG=inferior frontal gyrus; VS=vegetative state; MCS=minimally conscious state; EMCS=emergent from a minimally conscious state; LIS=Locked-In Syndrome; **=$p<0.01$; ***=$p<0.001$ (** and *** apply to power spectra only, as marked).

*Note.* aPatient MCS1 scored in the VS range immediately prior to his participation in the EEG assessment. However, this patient scored in the MCS Minus range in another CRS assessment several hours prior to his participation in this EEG investigation. For this reason, this patient has been classified as in a MCS Minus.