Electrophysiologic Evaluation of Diffuse Axonal Injury after Traumatic Brain Injury

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Abstract

Background: Electroencephalography may provide useful information about consciousness and cognitive processing in patients who have traumatic brain injury. Sleep disturbance after traumatic brain injury may impair cognitive function and affect rehabilitation.

Methods: In 15 patients who had chronic diffuse axonal injury after traumatic brain injury, electroencephalography and neuropsychologic tests were performed. In 8 patients who had subacute and chronic diffuse axonal injury and 7 healthy control subjects, electroencephalography, magnetoencephalography, and neuropsychologic tests were performed to evaluate sleep spindles and cognitive function.

Results: In the chronic stage of diffuse axonal injury, the mean peak frequency of alpha activity was significantly slower in patients who had abnormal than normal electroencephalography. The mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions were significantly slower in patients who had subacute diffuse axonal injury than healthy participants, and these parameters increased from the subacute to the chronic stage of diffuse axonal injury. After neurocognitive rehabilitation, cognitive functions were improved in all patients.

Conclusion: Alpha activity reflects the severity of disturbed consciousness in the acute stage after traumatic brain injury. Spindles will be an indicator of the recovery of consciousness in the chronic stage. Electroencephalographic makers may be useful in the diagnosis and prognosis of traumatic brain injury.

Keywords: Electroencephalography; Magnetoencephalography; Neuropsychologic; Cognition; Sleep spindles; Alpha activity

Abbreviations: DAI: Diffuse Axonal Injury; EEG: Electroencephalography; F: Female; GCS: Glasgow Coma Scale; GOS: Glasgow Outcome Scale; HDS-R: Hasegawa Dementia Scale Revised; M: Male; MEG: Magnetoencephalography; MMSE: Mini Mental State Examination; NA: Not available; PIQ: Performance Intelligence Quotient; TIQ: Total Intelligence Quotient; TBI: Traumatic Brain Injury; VIQ: Verbal Intelligence Quotient; WAIS-R: Wechsler Adult Intelligence Scale Revised; WISC-R: Wechsler Intelligence Scale for Children Revised

Introduction

In patients who have traumatic brain injury (TBI), a diagnosis and treatment are essential for acute patient care and long-term rehabilitation. Many problems may occur after TBI, including somatic problems (headache, fatigue, sexual dysfunction, and sleep disturbance), cognitive dysfunction (impairment of memory, attention, and information processing capacity) [1], mood changes (depression, aggression, emotional liability, and anxiety), and post-traumatic personality changes (self-centered behavior, reduced social awareness, and disinhibited emotions resulting from frontal and temporal lobe damage) [2,3].

Electroencephalography (EEG) has been evaluated for the clinical assessment of consciousness to support the diagnosis and prognosis [4,5]. Electrical activity of brain tissue may have good prognostic value after brain injury. When performed from 15 days to 4 years after injury, EEG may provide an objective and quantitative measure of the severity of brain injury [6]. Furthermore, EEG can detect early seizure activity, and provide information about sleep patterns during polysomnography. Sleep disturbances are common after TBI, and may include, insomnia, hypersomnia, and altered sleep-wake cycles [7-9]. However, there are no reproducible findings about specific changes in sleep quality and sleep architecture measured during polysomnography. Changes in sleep architecture after TBI are inconsistent, because the neural mechanisms contributing to sleep disturbances are multifaceted. The degree of damage to sleep-regulating centers such as the ascending reticular formation and associated pathways or neurotransmitter systems affect sleep disturbances [10,11]. Anxiety and depression frequently occur after TBI, and increased depression is associated with decreased sleep quality [12,13].

Proper diagnosis and effective treatment for sleep disorders after TBI, such as post-traumatic sleep-wake disturbances, possibly will improve the quality of sleep and improve the learning of motor tasks and cognition during sleep. Therefore, this study performed a study to evaluate the diagnosis utility of EEG in patients who had chronic diffuse axonal injury (DAI). In addition, simultaneously performed EEG and magnetoencephalography (MEG) in patients who had subacute to chronic DAI to evaluate the possibility of spindles as a biomarker.

Materials and Methods

Electroencephalography in the diagnosis of chronic diffuse axonal injury

The present study was approved by the ethics committee of National...
Rehabilitation Center for Persons with Disabilities. The study of EEG included 15 patients (13 male and 2 female; mean age, 23) who were diagnosed with acute DAI and who participated in our cognitive inpatient rehabilitation program. Coma duration was >6 hours, and magnetic resonance imaging showed little evidence of focal cerebral lesions.

The EEG recordings were performed in the chronic stage (mean time after injury 332±550 d), with 19 electrodes according to the international 10-20 system of electrode placement, and a linked ear reference (band-pass filter, 0.3 to 120 Hz, time constant, 0.3). The EEG data were evaluated initially by visual inspection, and fast Fourier transform was computed to calculate the power spectrum of alpha activity for each interval of 0.25Hz with a prism power spectrum (GE Marqettes, Tokyo, Japan). Ten epochs of stable alpha activity were summed and averaged. Cognitive functions were evaluated with neuropsychologic tests including Wechsler Adult Intelligence Scale- Revised (WAIS-R), which included Total Intelligence Quotient (TIQ), Verbal Intelligence Quotient (VIQ), and Performance Intelligence Quotient (PIQ); Mini-mental scale examination (MMSE); and Hasegawa Dementia Scale Revised (HDS-R).

Sleep spindles of patients who had subacute to chronic diffuse axonal injury

The study of sleep spindles included patients who had DAI (7 male and 1 female; age range 21 to 37 y; mean age 26 y) and 7 age-matched healthy volunteer control subjects (5 male and 2 female; age range, 24 to 32 y; mean age; 28 y).

Simultaneous EEG and MEG recordings were performed twice in the patients who had DAI: (1) during the sub-acute stage (mean time after injury 80 d), upon admission to the hospital for cognitive neurologic rehabilitation, and (2) during the chronic stage (mean time after injury, 151 d), upon completion of inpatient rehabilitation. In control subjects, the simultaneous EEG and MEG recordings were performed once.

Data were recorded during natural sleep (approximately 30 min) for each participant. The EEG recordings were made with a 60-channel electrode cap (Elekta-Neuromag, Finland) that was placed on the scalp according to the international 10-10 system of electrode placement. Magnetic signals were recorded from 204 planar gradiometers (Vectorview, Elekta-Neuromag, Finland) with a band-pass of 0.03 to 300 Hz for both the EEG and MEG signals. The sampling rate for digital conversion was 1001 Hz. For analysis, 10-stable fast spindles were selected by frequency and distribution. The Wechsler Adult Intelligence Scale III (WAIS-III) was used to evaluate cognitive functions.

Data analysis

Comparisons were performed with t test. Statistical significance was defined by P<0.05.

Results

Electroencephalography in the diagnosis of chronic diffuse axonal injury

All patients were diagnosed as DAI and were evaluated with EEG (Table1). In the chronic stage, 9 patients (60%) had abnormal EEG findings (Table2), including paroxysmal discharge noted in 4 patients (26%), centrotemporal spikes, 5 to 6 Hz small spikes and wave phantom. Cognitive function which was measured with WAIS-R, WAIS-III, MMSE, or HDS-R was similar in patients who had normal or abnormal EEG (Table 2). The mean peak frequency of alpha activity was significantly slower in patients who had abnormal than normal EEG (Table 2).

Sleep spindles of patients who had subacute to chronic diffuse axonal injury

Eight patients who had DAI and who were evaluated for sleep

| Patient No. | Age (y) | Sex | Acute stage | Chronic stage | Cognitive Function |
|-------------|---------|-----|-------------|----------------|--------------------|
|             |         |     | GCS | Coma Duration (d) | Time After Injury When EEG Performed (d) | MMSE | HDS-R | WAIS-R |
| 1           | 20      | M   | 7   | 4              | 150              | 29   | 29    | 82    | 91    | 75    |
| 2†          | 32      | M   | 7   | 7              | 120              | 23   | 22    | 48    | 60    | 45    |
| 3†          | 22      | M   | 7   | 30             | 210              | 27   | 26    | 96    | NA    | NA    |
| 4           | 14      | M   | 7   | 7              | 90               | 30   | 26    | 83    | 91    | 78    |
| 5           | 21      | M   | 5   | 15             | 90               | 23   | 19    | 58    | 66    | 60    |
| 6           | 17      | M   | 7   | 3              | 90               | 15   | 18    | 50    | 64    | 45    |
| 7           | 27      | M   | 7   | 100            | 300              | Tanaka-Binet test: 2 years | 10 | Scale out |
| 8           | 23      | M   | 7   | 10             | 150              | 27   | 28    | 64    | 71    | 67    |
| 9           | 27      | M   | 7   | 180            | 300              | Tanaka-Binet test: 2 years | Scale out |
| 10          | 20      | M   | 7   | 40             | 300              | 29   | 29    | 50    | 71    | 46    |
| 11          | 17      | M   | 7   | 30             | 240              | Tanaka-Binet test: 2 years | 8 | Scale out |
| 12          | 38      | F   | 7   | 14             | 180              | 23   | 28    | 84    | 82    | 89    |
| 13†         | 25      | F   | 9   | 80             | 2370             | 23   | 28    | 64    | 88    | 45    |
| 14†         | 26      | M   | 7   | 60             | 300              | 19   | 19    | 58    | 52    | 58    |
| 15†         | 17      | M   | 6   | 7              | 90               | 19   | 28    | 78    | 84    | 75    |
| Mean ± SD   | 23 ± 6  | 6.9 ± 0.8 | 39 ± 49 | 332 ± 570 | 24 ± 5 | 23 ± 7 | 68 ± 16 | 75 ± 13 | 62 ± 16 |

* N = 15 patients (13 male and 2 female). All patients had diffuse axonal injury. Abbreviations: EEG, electroencephalography; F, female; GCS, Glasgow Coma Scale; HDS-R, Hasegawa Dementia Scale Revised; M, male; MMSE, Mini Mental State Examination; NA, not available; PIQ, Performance Intelligence Quotient; TIQ, Total Intelligence Quotient; VIQ, Verbal Intelligence Quotient; WAIS-R, Wechsler Adult Intelligence Scale Revised; WISC-R, Wechsler Intelligence Scale for Children Revised; Scale out; could not perform neuropsychologic test due to severe mental dysfunction.
† Patient had subarachnoid hemorrhage (total, 5 patients)

Table 1: Clinical and cognitive characteristics of patients who had diffuse axonal injury after traumatic brain injury*.
spindles were young adult males who had moderate to severe injury (Table 3). In both healthy subjects and patients who had DAI, the centers of cortical activation of the 2 types of spindles (fast spindles and slow spindles) were located in 4 areas: the precentral areas of the right and left posterior frontal cortex and the postcentral areas of the right and left posterior parietal cortex. The precentral activation areas for fast spindles were located in the primary motor cortex and for slow spindles were located in the anterior frontal cortex, including the supplementary motor cortex. The postcentral activation areas for both types of spindles were located in the posterior parietal cortex. The postcentral activation areas included the parietal cortex of the primary sensorimotor areas and the posterior parietal areas. In healthy subjects, the cortical activation strength was similar between the four areas, including precentral and postcentral areas of both hemispheres (Table 4). The mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions were significantly slower in patients who had DAI, the mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions were significantly faster in the chronic than subacute stage (Table 4). The mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions were similar between patients who had DAI (chronic stage) and healthy participants (Table 4). There were no differences in mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions between hemispheres in healthy participants or patients who had DAI (subacute or chronic stages).

All 8 patients who had DAI had cognitive dysfunction, including memory disturbances and attention deficit. After neurocognitive rehabilitation, cognitive function improved in all patients, especially significant improvement in the WAIS-III subset scores for verbal intelligence quotient, performance intelligence quotient, total intelligence quotient, perceptual organization, and working memory (Table 5). All patients had favorable clinical outcome at 1 year after injury, with Glasgow Outcome Scale scores from 4 to 5. Some cognitive dysfunctions persisted after rehabilitation, but all patients returned home or resumed work.

**Discussion**

In the present study, the spontaneous EEG activity and cognitive function were evaluated after DAI. The EEG activity may be a marker of the severity of consciousness disturbance (acute stage) and recovery (chronic stage) after DAI. Although quantitative analysis was required for this study, alpha activity and spindles may be distinguished by visual inspection during typical clinical EEG studies [14]. Therefore, the present findings may be applicable to clinical practice. In patients who had TBI, it is important to consider the combined interaction and coordination of disturbed brain processes that are represented by the EEG, Glasgow Coma Scale, loss of consciousness, and neuropsychologic tests [15].

Previous quantitative EEG studies have attempted to develop an objective and quantitative measure of the severity of brain injury by using EEG obtained from 15 days to 4 years after injury [4,6,16]. The multivariate clinical assessment is a global variable that considers several neurophysiologic processes that are affected by TBI, especially those involving the frontal and temporal lobes [6].

The parieto-occipital alpha rhythm was attenuated by eye opening, visual stimuli, and by increased attentiveness [17]. These findings contributed to the hypothesis that oscillations may function as an 'idling' rhythm that characterizes an alert, yet inactive brain state. Recently published data have confirmed that alpha oscillations are strengthened by internal tasks, such as mental calculation and mental imagery [18-20]. The alpha band amplitude is increased during
the period of retention of short-term- and working memory and is suppressed subsequently. A previous study suggested that these large-amplitude oscillations during memory retention may inhibit the retrieval of memorized items, and this may be reflected in subsequent amplitude suppression. The present study showed longer coma duration in the acute stage, and slower frequency of alpha activity in the chronic stage of DAI. Further study may clarify whether alpha oscillations are associated with idling, inhibition, attention, or binding within the Global neural networks after brain damage that may cause consciousness disturbance in patients who have DAI.

In patients who have coma, the arousal system of the brain is deregulated because of diffuse brain or focal brainstem damage. In

| Characteristic | Healthy Subjects | Patients Who Had Diffuse Axonal Injury |
|---------------|------------------|----------------------------------------|
|               | Subacute         | Chronic                                |
| Peak frequency in central areas |                  |                                        |
| EEG (Hz)      |                  |                                        |
| Left          | 13.9 ± 0.2       | 13.4 ± 0.1*                            |
| Right         | 13.9 ± 0.2       | 13.5 ± 0.1*                            |
| MEG (Hz)      |                  |                                        |
| Left          | 13.7 ± 0.2       | 13.3 ± 0.1*                            |
| Right         | 13.7 ± 0.1       | 13.3 ± 0.1*                            |
| Amplitude in central areas |            |                                        |
| EEG (μV)      |                  |                                        |
| Left          | 29 ± 10          | 28 ± 10*                               |
| Right         | 30 ± 9           | 29.5 ± 0.1*                            |
| MEG (fT/cm)   |                  |                                        |
| Left          | 84 ± 29          | 68 ± 32*                               |
| Right         | 90 ± 34          | 70 ± 31*                               |
| Cortical activation source strength on MEG |                     |                                        |
| Precentral    |                  |                                        |
| Left          | 39 ± 19          | 22 ± 12*                               |
| Right         | 39 ± 20          | 17 ± 16*                               |
| Postcentral   |                  |                                        |
| Left          | 38 ± 21          | 20 ± 18*                               |
| Right         | 35 ± 17          | 21 ± 19*                               |

*N = 7 healthy subjects and 8 patients who had diffuse axonal injury. Data reported as mean ± SD. There were no differences in mean peak frequency of fast spindles, amplitude, and cortical activation source strength in precentral and postcentral regions between patients who had diffuse axonal injury (chronic stage) and healthy participants. Abbreviations: EEG, electroencephalography; MEG, magnetoencephalography
†Difference between healthy subjects and patients who had diffuse axonal injury (subacute stage).
‡Patients who had diffuse axonal injury; difference between subacute and chronic stages.

Table 4: Characteristics of fast spindles in healthy subjects and patients who had diffuse axonal injury after traumatic brain injury*

| Patient No. | Stage       | WAIS-III | VIQ  | PIQ  | TIQ  | Verbal comprehension | Perceptual organization | Working memory | Processing speed |
|-------------|-------------|----------|------|------|------|-----------------------|-------------------------|----------------|----------------|
|             |             |          |      |      |      |                       |                         |                |                |
| 1           | Subacute    | 76       | 63   | 65   | NA   | NA                    | NA                      | NA             | NA             |
|             | Chronic     | 102†     | 96†  | 98†  | NA   | NA                    | NA                      | NA             | NA             |
| 2           | Subacute    | 56       | 50   | 49   | 54   | 55                    | 54                      | 50             | 50             |
|             | Chronic     | 112†     | 98†  | 101† | 98   | 102†                  | 98†                     | 97             | 97             |
| 3           | Subacute    | 96       | 102  | 98   | 86   | 106                   | 103                     | 92             | 92             |
|             | Chronic     | 110†     | 98†  | 102† | 98   | 110†                  | 111†                    | 92             | 92             |
| 4           | Subacute    | 88       | 84   | 85   | 90   | 83                    | 100                     | 84             | 84             |
|             | Chronic     | 111†     | 97†  | 113† | 102  | 110†                  | 92                      | 102†           | 102†           |
| 5           | Subacute    | 59       | 58   | 55   | 61   | 61                    | 60                      | 57             | 57             |
|             | Chronic     | 86       | 72†  | 88   | 72   | 82†                   | 88†                     | 66             | 66             |
| 6           | Subacute    | 64       | 74   | 65   | 56   | 66                    | 62                      | 63             | 63             |
|             | Chronic     | 84†      | 102† | 91†  | 92   | 119†                  | 76†                     | 78             | 78             |
| 7           | Subacute    | 50       | 53   | 103  | 86   | 112                   | 111                     | 60             | 60             |
|             | Chronic     | 90†      | 103† | 95†  | 86   | 112†                  | 111†                    | 60             | 60             |
| 8           | Subacute    | 114      | 90   | 104  | 109  | 101                   | 111                     | 81             | 81             |
|             | Chronic     | 120      | 94   | 110  | 110  | 118†                  | 110                     | 92             | 92             |

*N = 8 patients. Abbreviations: PIQ, Performance Intelligence Quotient; TIQ, Total Intelligence Quotient; VIQ, Verbal Intelligence Quotient; WAIS-III, Wechsler Adult Intelligence Scale III. NA: not available
†Difference between subacute and chronic stages: P ≤ .05

Table 5: Cognitive function in patients who had traumatic brain injury*
coma the EEG may have various patterns of activity including, a
generalized slowing in the delta or theta range, alpha- coma, spindle-
coma, burst- suppression, and epileptiform activity [21-23]. The normal
sleep-wake cycle is disrupted or completely absent in coma. However,
the coma tracing resemble normal wakefulness or normal sleep [24].
Spindle coma, characterized by the occurrence of spindles in comatose
patients, may be caused by central nervous system trauma, infection,
or metabolic encephalopathy. Abnormal spindle formations occur
because of midbrain involvement with sparing of thalamic structures
[25]. In patients who have supratentorial lesions, spindle coma suggests
the presence of intact cortical function and good prognosis [26,27].
Spindle- coma is considered as a benign form of coma, with EEG
reactivity to stimuli that may herald a favorable outcome [28,29].
In comatose patients, spindles are best demonstrated during first few
days after trauma. In a previous study, spindle activity was observed in 91%
patients who had posttraumatic coma, and 30% patients developed
prolonged coma [29].

In the acute stage of coma, spindles may have prognostic value.
Symmetric occurrence of spindles may be associated with a good
prognosis, and asymmetric or decreased spindles may be associated
with a poor prognosis [26]. In patients who have chronic coma,
recovery may occur over many months or years. Focal injury patterns
in the central thalamus are associated with coma, vegetative state,
and a minimally conscious state. Medial components of the posterior
intralaminar region of the central thalamus are involved in spindle
generation. Therefore, the recovery of consciousness may be strongly
associated with the recovery of sleep spindles after brain injury [30,31].

Sleep disorders are common after the acute phase of TBI. Cognitive
dysfunctions may be caused by sleep disturbances and may worsen
the impairment in patients who have TBI. Sleep disturbances may
include hypersomnia, insomnia, altered sleep-wake cycles, periodic
limb movements in sleep, and disorders of rapid eye movement sleep.
Breathing disorders of rapid eye movement sleep such as obstructive
sleep apnea or central apnea. In patients who have obstructive sleep
apnea have greater impairment of neurocognitive function, especially
memory and sustained attention, than patients who do not have disordered
breathing during sleep.

In patients who have TBI, findings of sleep architecture are
inconsistent and may include; no changes, increased slow waves,
decreased rapid eye movement sleep, increased rapid eye movement
sleep during the second half of the night, no change in rapid eye
movement sleep, or decreased onset latency of rapid eye movement
sleep. Sleep-wake regulating centers and associated pathways are
damaged in TBI, and these damages will be the cause of disturbances
of sleep architecture. Sleep disturbances contribute to fatigue, which may
be associated with mental slowness and, slowed information processing
[32]. Sleep disturbances also contribute to anxiety and depression.
Patients with TBI may have disrupted circadian regulation of melatonin
synthesis, including lower levels of melatonin production during the
evening [12]. Reduced sleep quality induces depression. Early diagnosis
and treatment with modafinil, melatonin, and light therapy may
improve alertness and mood. It also is important to support lifestyle
modifications, and facilitate daytime activities [33].

In summary, the present study showed that electrophysiologic
markers such as alpha activity and sleep spindles may be useful in the
diagnosis and prognosis of TBI. The multivariate clinical assessment is
a global variable that may combine several neurophysiologic that are
needed to access prognosis.

Proper diagnosis and effective treatment of sleep disorders after
TBI may potentially improve the quality of sleep, the off-line learning
of motor tasks and cognition during sleep, alertness, mood, and
rehabilitation.

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