Abstract

The identification of the predictive factors and biological markers associated with treatment-related changes in the symptoms of Internet gaming disorder (IGD) may provide a better understanding of the pathophysiology underlying this condition. Thus, the present study aimed to identify neurophysiological markers associated with symptom changes in IGD patients and to identify factors that may predict symptom improvements following outpatient treatment with pharmacotherapy. The present study included 20 IGD patients (mean age: 22.71 ± 5.47 years) and 29 healthy control subjects (mean age: 23.97 ± 4.36 years); all IGD patients completed a 6-month outpatient management program that included pharmacotherapy with selective serotonin reuptake inhibitors. Resting-state electroencephalography scans were acquired prior to and after treatment, and the primary treatment outcome was changes in scores on Young’s Internet Addiction Test (IAT) from pre- to posttreatment. IGD patients showed increased resting-state electroencephalography activity in the delta and theta bands at baseline, but the increased delta band activity was normalized after 6 months of treatment and was significantly correlated with improvements in IGD symptoms. Additionally, higher absolute theta activity at baseline predicted a greater possibility of improvement in addiction symptoms following treatment, even after adjusting for the effects of depressive or anxiety symptoms. The present findings demonstrated that increased slow-wave activity represented a state neurophysiological marker in IGD patients and suggested that increased theta activity at baseline may be a favorable prognostic marker for this population.

Abbreviations: BAI = Beck anxiety inventory, BDI = Beck depression inventory, IAT = Young’s Internet Addiction test, IGD = Internet gaming disorder, IQ = intelligence quotient.

Keywords: Internet gaming disorder, resting-state EEG, slow-wave activity, treatment response

1. Introduction

Internet gaming disorder (IGD) refers to compulsive and problematic use of Internet-based games that interferes with normal daily life.[1] Previous research has shown that various psychiatric comorbidities are associated with IGD, including depressed mood,[2] anxiety,[3] attention deficit hyperactivity disorder,[4] and obsessive–compulsive disorder.[5] Electroencephalographic (EEG) studies of spontaneous brain activity during resting conditions with the eyes closed have been increasingly recognized as a useful method for the investigation of the neural correlates of cognition and behavior.[6] Spontaneous brain activity is neural activation in the absence of an explicit task, such as sensory input or motor output, and, hence, is referred to as resting-state activity.[7] This type of neural activity reflects the brain’s ability to allocate resources and prepare for changes in the internal and external environments.[8] Thus, the examination of resting-state EEG data as a measure of cognitive capabilities may enhance the current understanding of basic brain functions.

Several studies have used resting-state EEG to investigate the characteristics of patients with IGD. Absolute beta power is lower in these patients under resting-state conditions and is correlated with standard self-report measures of impulsivity.[9] Furthermore, the resting-state EEG activity of IGD patients aids in the differentiation of this population from patients with alcohol use disorder (AUD) because IGD patients have lower absolute beta power than AUD patients and healthy controls. However, this marker is not related to the severity of IGD, which suggests that...
decreased beta power is a trait marker. In contrast, AUD patients exhibit higher absolute delta power than do patients with IGD and healthy controls.IGD patients also frequently exhibit increased relative theta, and decreased relative alpha powers in all regions, especially fronto-central regions, which are known to be closely related to depression. Therefore, it is necessary to diagnose the individual comorbid psychiatric symptoms of IGD patients to accurately characterize the neurobiological characteristics associated with this disorder.

The identification of biological markers associated with treatment-related changes in IGD symptoms can provide a better understanding of the pathophysiology underlying IGD. Because IGD is associated with a poor quality of life and lowered psychological well-being, the early identification of treatment responders and the implementation of early interventions involving individualized treatment approaches would likely have significant clinical importance. Abstaining AUD patients show an increase in slow-wave brain activity after 6 months of treatment compared with baseline measurements, which indicates a normalization of brain function. Recently, Wang et al. suggested that abnormal resting-state electrical activity, including increased power in the theta band, in the brains of patients who use illicit opiates may be reduced by methadone treatment. Moreover, an event-related potential study found that the P300 amplitude reliably predicts treatment completion in substance-dependent populations; however, this peak was not recorded during the resting state.

To date, no studies have investigated the neurophysiological factors associated with treatment responses in patients with IGD. Thus, the present study aimed to determine the neurophysiological markers associated with symptom changes in IGD patients with comorbid psychiatric issues and to detect markers that will predict symptom improvements following outpatient management with pharmacotherapy. Based on previous studies of patients with IGD and/or substance use disorders, it was hypothesized that IGD patients with comorbid symptoms would exhibit increased activities in the delta and theta bands on the fronto-central regions and also alterations between slow-wave activity in the delta and theta bands and improvements in addiction symptoms following 6 months of outpatient management.

2. Methods and materials

2.1. Study participants

The present prospective longitudinal study included 49 male participants, 20 of whom were IGD patients (mean age: 22.71 ± 5.47 years) and 29 of whom were healthy control subjects (mean age: 23.97 ± 4.36 years). All patients had sought treatment at an outpatient clinic of SMG-SNU Boramae Medical Center in Seoul, South Korea due to excessive participation in Internet gaming, and IGD was diagnosed by an experienced psychiatrist based on the criteria of the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5). Additionally, Young’s Internet Addiction Test (IAT) was administered to assess the severity of IGD symptomatology. The IAT has been extensively used in the research field of IGD and well documented its psychometric properties in different population and languages. A recent validation study for Korean translation of the IAT showed outstanding internal consistency and high test–retest reliability. These items are rated on a 5-point scale ranging from 1 (very rarely) to 5 (very frequently); in the present study, IAT scores were calculated with the total score for all 20 items ranging from 20 to 100. The present study included only those patients with IAT scores ≥70 who spent more than 4 hours per day and 30 hours per week using Internet games to clarify the pathological changes associated with IGD and also to evaluate only patients with severe IGD rather than patients at a high risk of developing this disorder. Additionally, to assess lifetime psychiatric diagnoses, the Structured Clinical Interview for DSM-IV disorders was used.

Following the completion of the clinical assessments and a baseline EEG scan, the IGD patients completed a 6-month outpatient management program that included pharmacotherapy with selective serotonin reuptake inhibitors using the average following doses: escitalopram at 15.83 ± 9.17 mg, fluoxetine at 50.00 ± 9.17 mg, and paroxetine at 30.00 ± 14.14 mg. All IGD patients who completed the 6-month treatment program had comorbid depressive or anxiety symptoms at baseline and completed a follow-up EEG scan upon finishing. The primary treatment outcome of the present study was change in IAT score from pre- to posttreatment, and the Beck Depression Inventory (BDI) and Beck Anxiety Inventory (BAI) were administered at the pre- and posttreatment assessments of the IGD patients to evaluate changes in comorbid depressive and anxiety symptoms. Healthy control subjects were directly recruited from the local community, did not have a history of psychiatric disorders, and played Internet games less than 2 hours per day.

Participants were excluded from the present study if they had a history of significant head injury, seizure disorder, intellectual disability, psychotic disorder, or substance use disorder (other than one involving nicotine). Additionally, all participants were medication-naïve at the time of the baseline assessment. The Korean version of the Wechsler Adult Intelligence Scale–III was administered to all subjects to estimate their IQ; only individuals with Wechsler Adult Intelligence Scale–III scores ≥80 were included in the present study. The Institutional Review Board of the SMG-SNU Boramae Medical Center approved the study protocol, and all subjects provided written informed consent prior to participation.

2.2. EEG recording

The detailed protocols for the EEG recording and data acquisition procedures have been described in previous reports published by our research group. Brieﬂy, participants were engaged in a resting state, while seated in a darkened noiseless room that was connected to a recording room via a 1-way glass window. The EEG recording was conducted for 10 minutes under the following conditions: 4 minutes with eyes closed, 2 minutes with eyes open, and 4 minutes with eyes closed.

The EEG recordings and acquisitions were obtained using SynAmps2 with a 64-channel Quick-cap and the NeuroScan system (Scan 4.3, Compumedics Ltd; Abbotsford, Australia). A single channel with bipolar electrodes was attached to the mastoids as a reference, and a ground channel was placed between the Fpz and Fz electrodes. The EEG signals were obtained at a sampling frequency of 500 Hz and band-pass filtered at 0.1 to 60 Hz using Scan 4.3 with an electrode impedance below 5 KΩ. The recordings from the NeuroScan system were transferred to NeuroGuide software (NG 2.5.5, Applied Neuroscience, Inc, St. Petersburg) for spectral analysis in
a 32-bit file format, and 19 of the 64 channels were driven by the following NeuroGuide montage set: FP1, F3, F7, Fz, FP2, F4, F8, T3, C3, Cz, T4, C4, T5, P3, O1, Pz, T6, P4, and O2.

Artifact removal was performed off-line using the artifact rejection toolbox in the NeuroGuide software. Additionally, all EEG recordings were visually analyzed to eliminate signal distortions, such as eye muscle movements. Samples were selected by visual inspection to obtain a minimum of 20 to 60 seconds of data during the eyes-closed resting state for the spectral analysis, and the accepted epochs of EEG data for both the absolute (actual spectral power; $\mu V^2$) and relative proportion of spectral power in the combined sum of frequency bands; %) powers were smoothed using fast Fourier transforms. The power was divided into 4 frequency bands and averaged with the NeuroGuide spectral analysis system as follows: delta (0.5–4 Hz), theta (4–8 Hz), alpha (8–12 Hz), and beta (12–30 Hz). In the analysis of the present study, we focused on the delta and theta bands of slow-wave. Additionally, based on previous studies, we analyzed EEG activity from 19 selected sites that were divided into 3 regions by averaging within each region: frontal (FP1, F3, F7, Fz, FP2, F4, and F8), central (T3, C3, Cz, T4, and C4), and posterior (T5, P3, O1, Pz, T6, P4, and O2).[9–11]

2.3. Statistical analysis

Prior to the formal analyses, an exploratory data analysis was conducted to identify and remove outliers to avoid the possibility of spurious results. Because repeated or multiple outcomes from the same subject are correlated, the present study utilized a generalized estimating equation, which is an extension of the generalized linear model for multivariate responses, to assess group effects on the absolute and relative powers in each delta and theta frequency band. The generalized estimating equation was employed to test group (IGD vs. healthy control), region, and the interaction of these factors in each frequency band because this model has previously been used to analyze EEG characteristics.[19–21] Significant interactions between the groups in different regions were calculated using a group-by-region interaction because it reflects the group effects on the absolute and relative powers for each frequency band according to region. In the absence of an interaction effect, the group effects were also tested.

Independent-sample $t$ tests were performed to compare the demographic and clinical variables of the 2 groups, whereas differences in clinical symptoms and EEG data between the pre- and posttreatment assessments were analyzed with a repeated measure analysis of variance after adjusting for BDI and BAI scores. Additionally, a multiple linear regression analysis was conducted to examine associations among the baseline EEG markers and changes in IAT scores (IAT score at pretreatment − IAT score at posttreatment) after adjusting for BDI and BAI scores. A Pearson correlation analysis was performed to examine relationships among changes in EEG markers and changes in IAT scores after 6 months of treatment. All statistical analyses were performed with the SPSS Statistics package, version 20 (IBM Inc; NY) and R version 2.15.2 (http://www.r-project.org). $P$ values $<0.05$ were considered to indicate statistical significance.

3. Results

3.1. Demographic and clinical/cognitive data

Table 1 summarizes the demographic and clinical characteristics of the participants. There were no significant differences between the IGD patients and healthy control subjects in terms of age, education, or IQ, but the IGD patients had higher BDI ($P<0.001$) and BAI ($P<0.001$) scores than the controls. After 6 months of treatment, the IGD patients exhibited a significant decrease in IAT scores ($P<0.001$) but not in BDI and BAI scores.

3.2. Quantitative changes in EEG data following treatment

Figure 1 illustrates the scalp topographies of patients in the IGD group before and after treatment in terms of the absolute and relative powers in each band. Compared with the healthy control group, the IGD group had increased absolute powers in the delta band of the total brain (estimate $=4.96$, $t=2.05$, $P=0.046$) and in the theta band of the central brain region (estimate $=5.15$, $t=2.41$, $P=0.021$; Fig. 2). There were no group-by-region interaction effects for the absolute or relative powers in any of the other bands, and the main group effects were not significant in any of the regions.

Following 6 months of treatment, the absolute power in the delta band of the frontal region of the IGD group exhibited a significant

Table 1

| Demographic characteristics of study subjects. | IGD patients (n = 20) | Healthy controls (n = 29) | $t$ | $P$ value |
|---|---|---|---|---|
| **Demographic data** | | | | |
| Age, y | 22.71 (5.47) | 23.97 (4.36) | −1.00 | 0.318 |
| Education, y | 13.09 (2.42) | 14.19 (2.13) | −1.78 | 0.083 |
| IQ score | 114.06 (17.21) | 119.31 (9.49) | −1.46 | 0.152 |
| **Clinical data** | | | | |
| Duration of illness, y | 7.32 (3.68) | – | – | – |
| IAT | 75.45 (7.35) | 27.04 (8.84) | 15.91 | $<0.001^*$ |
| BDI | 18.88 (10.90) | 4.15 (3.22) | 7.26 | $<0.001^*$ |
| BAI | 19.76 (15.67) | 6.88 (4.01) | 4.24 | $<0.001^*$ |

| Pretreatment (n = 20) | Post-treatment (n = 20) | $F$ | $P$ value |
|---|---|---|---|
| IAT | 75.45 (7.35) | 59.25 (20.81) | 15.90 | $<0.001$ |
| BDI | 18.88 (10.90) | 14.82 (12.87) | 0.22 | 0.653 |
| BAI | 19.76 (15.67) | 13.00 (12.26) | 0.88 | 0.364 |

Data are presented as means (standard deviations) and were analyzed with independent-sample $t$ tests and repeated measure ANOVAs.

ANOVA = analysis of variance, BAI = Beck anxiety inventory, BDI = Beck depression inventory, IAT = Internet Addiction Test, IQ = intelligence quotient.

* $P<0.05$. 


reduction compared with baseline \((z=2.03, P=0.043)\), and the extent of this decrease was significantly correlated with change in IAT score \((r=0.57, P=0.011)\). In contrast, there were no significant absolute or relative changes in any of the other bands following treatment nor were there significant differences in the absolute or relative powers in any region between the IGD group at the posttreatment assessment and the healthy control group. Further, the reduction of the frontal delta activity was not correlated with changes in depressive \((r=-0.08, P=0.778)\) or anxiety symptoms \((r=-0.02, P=0.935)\) following treatment in the IGD group.

**Figure 1.** Topographical maps of the absolute and relative powers in patients with Internet gaming disorder (IGD) before and after the 6-month outpatient treatment and the healthy control group at baseline. Scales show \(\mu V^2\) for absolute power and \% for relative power. Red represents higher values, and blue represents lower values.

**Figure 2.** Absolute and relative powers in each band under the resting-state eyes-closed condition. Horizontal bars represent standard deviations. * \(P<0.05\).
3.3. Quantitative EEG components predictive of improvements in IGD symptoms

A multiple regression analysis revealed that change in IAT score was significantly predicted by absolute power in the theta band of the central brain region at baseline (estimate $= 3.10, t = 2.53, P = 0.025$; Fig. 3). In other words, higher absolute theta activity in the central region at baseline predicted a greater possibility of improved addiction symptoms in IGD patients with comorbid issues following treatment, even though the effects of depressive or anxiety symptoms at baseline were controlled.

4. Discussion

The present study was the first to investigate the neurophysiological changes associated with symptom improvement in IGD patients. IGD patients with comorbid depressive or anxiety symptoms showed increased resting-state EEG activity in the delta and theta bands at baseline, but this increased delta band activity was normalized after 6 months of pharmacotherapy. Additionally, higher absolute theta activity at baseline predicted a greater possibility of improved addiction symptoms following treatment, but these neurophysiological changes were not related to depressive or anxiety symptoms because depressive or anxiety symptoms did not show significant changes following treatment; there were no correlations between the reduction of the frontal delta band activity and depressive or anxiety symptoms changes following treatment; and higher absolute theta activity in the central region at baseline predicted a greater possibility of improved addiction symptoms in IGD patients with comorbid issues following treatment, even though the effects of depressive or anxiety symptoms at baseline were controlled. The present findings indicated that slow-wave activity in IGD patients may be a state marker associated with changes in addiction symptoms following treatment.

Resting-state EEG activity represents the readiness or potential of the brain to allocate neural resources.[27] The default mode network is a network of brain regions that exhibits increased activity during the resting state, and the intrinsic activities within these areas are known to impact subsequent stimulus or task-induced activity.[28] Therefore, investigating resting-state brain activity may further the current understanding of basic brain functions. Delta and theta wave activities are involved in a variety of cognitive processes. For example, increased theta power is observed during tasks that demand attention, which suggests that low-frequency bands are involved in higher-order control processes.[29] Moreover, event-related P300 responses are primarily the outcome of delta and theta oscillations elicited during the cognitive processing of stimuli.[30,31] These P300-related oscillations are associated with different cognitive functions. For example, theta is associated with memory processes or attention and is considered to be an index of frontal processing, whereas delta is related to signal detection and decision-making and is generated by cortico-cortical interactions.[32] Patients with AUD exhibit significantly lower evoked delta and theta oscillation amplitudes during the processing of target stimuli.[33] Similarly, a recent study reported that the auditory P300 amplitude is reduced in patients with IGD compared with healthy controls.[34] Given that resting-state brain activity impacts subsequent task-induced activity, it is of interest to note that increased delta and theta activities during the resting state may be related to decreased task-related P300 amplitudes in IGD patients. Additionally, EEG activity in the slow frequency bands may be associated with inhibitory control. For example, Schiller et al.[35] found that increased power in the slow frequency bands of the lateral prefrontal cortex is linked to inhibitory deficits and, moreover, impairments in inhibitory function may occur in patients with IGD. Therefore, increased delta and theta activities during the resting state are likely associated with dysfunctional inhibitory control in IGD patients.

In the present study, IGD patients exhibited significant improvements in their addiction symptoms. Additionally, the increased delta band activity observed at baseline in this group was normalized after treatment and was significantly correlated with improvements in IGD symptoms. Moreover, the present study found that higher absolute theta activity at baseline predicted a greater possibility of improvement in IGD symptoms after treatment. After controlling for the effects of depressive or anxiety symptoms, these findings remained significant. There have been mixed results regarding changes in slow-wave activity and their associations with treatment in AUD patients. Saletu-Zyhlarz et al.[12] reported that abstaining patients showed an increase in slow activity after 6 months of treatment compared with baseline, and Pollock et al.[36] found persistent slow-wave alterations in alcoholics with prolonged abstinence. These authors also suggested that the higher power of the slow waves could be considered a trait marker of AUD.[36] In contrast, the present study found that IGD patients displayed a decrease in absolute delta power; in other words, absolute delta power in this population normalized after 6 months of treatment and was associated with an improvement in IGD symptoms. This finding indicates that there may be different mechanisms underlying the neurophysiological activities associated with symptom changes in AUD and IGD patients. As mentioned above, slow-wave activities are involved in various types of cognitive processes.[26–28] Thus, the normalized delta power in IGD patients following treatment was likely related to changes in cognitive function. Accordingly, patients with Alzheimer disease exhibit significant decreases in power in the delta and theta bands during treatment with rivastigmine, which is an acetylcholinesterase

![Figure 3](https://example.com/figure3.png)

Figure 3. Relationship between absolute power in the theta band of the central region at the pretreatment assessment and change in IAT score after 6 months of treatment in patients with IGD. Higher absolute theta activity in the central region at baseline predicted a greater possibility of improvements in addiction symptoms after treatment in IGD patients with comorbid problems. IAT = Young Internet Addiction Test, IGD = Internet gaming disorder.
in clinical settings have various types of comorbid issues; thus, the present findings may provide useful clinical information regarding the neurobiological markers associated with IGD.

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