Introduction: Our aim in this study was to investigate spectral power density (PSD) in first-episode mania and subsequent remission period and to evaluate their difference.

Methods: Sixty-nine consecutive cases referring to our hospital within the previous 1 year, who were evaluated as bipolar disorder manic episode according to The Diagnostic and Statistical Manual of Mental Disorders-IV (DSM-IV) at the first episode and had the informed consent form signed by first degree relatives, were included in this study. Exclusion criteria included having previous depressive episode, using drugs which could influence electroencephalographic activity before electroencephalography (EEG), and having previous neurological disease, particularly epilepsy, head trauma, and/or loss of consciousness. EEG records were obtained using a digital device in 16 channels; 23 surface electrodes were placed according to the International 10–20 system. Spectral power density (dbµV/Hz) of EEG signal provided information on the power carried out by EEG waves in defined frequency range per unit frequency in the present study.

Results: A peak power value detected on the right with FP2P4 and on the left with F7T3 electrodes were found to be higher in the manic episode than in the remission period (p=0.018 and 0.025). In the remission period, in cases with psychotic symptoms during the manic period, F4C4 peak power value was found to be lower than that in cases with no psychotic findings during the manic period (p=0.027). There was no relation was found between YMRS scores and peak power scores.

Conclusion: Electrophysiological corollary of mood episode is present from the onset of the disease, and it differs between the manic and remission periods of bipolar disorder. In the remission period, peak power values of PSD distinguish cases with psychotic findings from cases without psychotic findings when they were manic.

Keywords: First episode, mania, QEEG, spectral power density, frontal cortex, temporal cortex

INTRODUCTION

In bipolar disorder (BD), cognitive impairment in various areas, particularly attention, verbal learning, and executive function, is observed during active disease and remission periods (1,2,3). Savitz et al. (4) have suggested that cognitive function in BD is the consequence of chronic impairment in the neural network. It has been demonstrated that frontolimbic connections are influenced in BD (5). However, for an integrated mood and cognitive function, corticocortical connections should also be intact. In addition, synchronization among neurons is also required.

This is made possible by electric signals (6,7). Electroencephalography (EEG) is one of the current most optimum means of investigating dynamic brain functions owing to its high temporal resolution. Quantitative electroencephalography analysis converts a certain EEG characteristics into a numerical value. A signal analysis method used commonly in quantitative EEG analysis is EEG power spectrum density (PSD) signal analysis, which yields information on the power carried by EEG waves per unit frequency at a defined frequency range (8).

In BD, El-Badri et al. (9) found in 2001 with 29 euthymic cases that 29 EEG PSD was higher than healthy controls in all bands. However, the greatest difference was found in left occipital beta band. The aim of this study is to investigate EEG PSD in first-episode mania and the subsequent remission periods, to determine whether they are different, and to investigate whether peak values of PSD and clinical characteristics are related.

METHODS

Sample

In the present study, 69 consecutive first-episode mania cases aging 18–65 whose informed consent was signed by first degree relatives and were referred to our outpatient clinic or emergency service within the previous year and diagnosed with BD manic episode, ac-
cording to The Diagnostic and Statistical Manual of Mental Disorders-IV (DSM-IV) were evaluated. No previous depressive period; no history of previous neurological disease, particularly epilepsy, head trauma, and/or loss of consciousness; and not using any drug (antiepileptics, anxiolytics, antidepressants, and/or antipsychotics) that could affect EEG activity before EEG recording were the inclusion criteria. Some cases were excluded as they had to be administered a pharmacological agent that may affect EEG before the recording of EEG (n=5), some because their diagnosis was changed to schizoaffective disorder or psychotic disorder (n=3), and some because they did not come for control EEG recording in the remission period (n=11).

Finally, data of overall 50 patients were considered for evaluation.

Among first BD manic episode cases, epileptic EEG was found in two cases (4%) and nonepileptic abnormal EEG in 10 cases (20.8%). Peak values of PSD were compared after excluding cases with epileptic or nonepileptic abnormal EEG findings (n=38).

**Tools**

**DSM-IV structured clinical interview (SCID-I):** It was developed by First et al. (10) in 1997 to investigate first axis disorders according to DSM-IV, and its reliability and validity study in Turkish was performed by Özkürkçügil et al. (11).

**Mood disorders diagnosis and follow up form (SKIP-TURK):** It was used to investigate sociodemographic and clinical characteristics of the cases (12).

**Young mania rating scale (YMRS):** It was used to determine the severity of manic symptoms before treatment and to confirm remission. This scale that was filled out by the interviewer was developed by Young et al. (13), and its validity and reliability study in Turkish was performed by Karadağ et al. (14).

**Implementation**

Necessary approval for the study was obtained from Erenköy Psychiatric and Neurological disorders Hospital by the same technician using a 33-channel ESAQATE, Mizar, 2004/Italy digital EEG device in 16 channels. Twenty-three surface electrodes were placed according to the International 10–20 system (8). EEG recordings were obtained at a 70-Hz sampling speed in the resting position, and an artifact control was created before the evaluations.

The following points were determined: (i) whether EEG was epileptic; (ii) if not epileptic, whether it included other abnormal patterns (dysrhythmia, slow wave, amplitude abnormalities, and asymmetry); (iii) peak power values for 16 channels at 12–30 Hz (alpha and beta band) frequency range. We chose this frequency range because it included both alpha and beta activity. Spectral analysis is a commonly used signal analysis method in EEG analysis. To observe the frequency corresponding to density, common approach in HFD analysis is using the squares of Fourier coefficients to form the graphic termed as power spectrum and the digital EEG device presents these data with a loaded program. PSD EEG signal provides information regarding the power carried by the EEG waves at the defined frequency range per unit frequency. In the present study, PSD was defined as dBµV/Hz., i.e., output power is logarithmic and expressed in decibels. Maximum value was considered among the power values at the given frequency range (12–30 Hz) and frequency corresponding to this value was found. This procedure was performed for each channel.

**Statistical Analysis**

For the comparison of numerical data, T test (mean±SD<0.3) was used; if they were not distributed normally Mann–Whitney U test was used. For the comparison of categorical data, chi-square and Fisher’s exact tests were used. All tests were two ended and p-value of <0.05 was considered significant. For correlation analysis, Pearson correlation test was used. For the comparison of repeated measurements in the same cases, Wilcoxon test was used.

**RESULTS**

**Sample Characteristics**

Of the BD cases in the first-episode manic period, 44% were female and 56% male. Their mean age was 26.4±4.8 years and mean duration of education was 10.7±2.1 years (Table 1).

**Comparison of Peak Power Values between Mania and Remission**

Peak power values detected at FP2P4 and F7T3 electrodes were found to be higher during the manic period than those during the remission period (p=0.027 and 0.033; Table 2). These values correspond to approximately 15- and 16-Hz in beta band.

**Investigation of the Relation between Peak Power Value and Clinical Characteristics**

**Manic period:** In first-episode mania cases, cases with significant family history, childhood trauma, suicide attempt, and psychotic findings were compared with the cases without them, and peak power values were found to be similar between both. No relation was found between age (also the age of onset) and YMRS scores and peak power values at 16 points.

**Remission period:** When peak power values were compared between cases with significant family history, childhood trauma, suicide attempt, and psychotic findings and those without them, similar to the manic period, peak power values were found to be similar. However, in cases with psychotic findings during the manic period, F4C4 peak value in the remission period was found to be higher than that in those without psychotic findings during the same period (p=0.027). These values correspond to approximately alpha band 11 Hz.

**DISCUSSION**

In the study by Ray and Cole (16) in 1985, it was reported that in cognitive functions such as mental arithmetic not requiring environmental attention, alpha activity increased, particularly in the parietal region, whereas in the 195
In ADHS, beta activity was found to be higher than that in healthy controls, particularly in the frontal region, and it was reported that it decreased with drugs but did not return to normal and was associated with hyperactivity and impulsivity (19). In ADHS, frontal beta activity is higher in cases responding to stimulants than those not responding (20). In addition, frontal beta activity was found to be associated with improvement in Conners Continuous Performance Test performance (maintenance of attention and inhibitor control) and decrease in hyperactivity–impulsivity scores reported by the family. Increased beta oscillatory beta activity after a positive feedback has been repeatedly demonstrated in previous studies (21). In gambling test including both positive and negative feedback, unexpected awards were markedly associated with increased beta activity (22).

In a study comparing cases who are successful in inhibitor control with those who are not so in stop signal task, highest response in intracranial electrodes was obtained in the right inferior frontal gyrus beta band (optimally approximately 16 Hz) (23). In the same study, in cases with inhibitor control, alpha/beta desynchronization in primary motor cortex was found to be decreased. It was suggested that this decrease is associated with GABAAergic inhibition. It was also proposed that the function of GABA in this context is a type of gating to synchronize and harmonize processes in different areas (24,25,26). In the study by Liu et al. (27), it was demonstrated that increased slow-wave activity in frontal central regions was specific to bipolar depressions. In a study on early onset psychosis cases, it was suggested that abnormal gama and beta activities detected with magnetoencephalography (MEG) implied reduction in control over motor movements, which was associated with the reflection of impaired inhibition function of GABA at cellular and system level (28). In the present study, no relation was found between YMRS scores and peak values at 16 points. However, the presence of psychotic finding appears to create a difference. F4C4 peak value during the remission period was found to be lower in cases with psychotic findings during manic period than those without psychotic cases during the same period. This finding may be interpreted as the trace left by psychosis. In the course of BD, disease periods have accumulating effect, which is felt most strongly in cognitive functions (23).

Central alpha activity, corresponding to peak power values, which are found to be lower in the remission period in cases with psychotic findings in the mania period, are associated with coding and recognition processes (29). Increase in performance in working memory tests, particularly semantic memory, is expected to transfer long term memory codes to short term memory and to reactivate them. In the study by Bajar et al. (30), alpha activity detected in euthymia was found to be lower in cases with BD than that in healthy controls. The relation between psychotic findings and EEG abnormality is more marked in psychotic spectrum. Increased gamma activity in first-attack psychotic cases is a finding corroborated in many studies (31). In cases in which EEG abnormality was detected in first attack, unfavorable clinical course characteristics were observed more commonly in the first 3-year follow-ups (32). In a study comparing epileptic cases with and without psychotic findings, in epileptic cases with schizophrenia-like psychosis, increased alpha synchronization was observed in MEG and dorsolateral prefrontal cortex (33).

The effect of psychotrophic drugs on peak values during the remission period is an important confounding factor. As in previous studies with similar design, the fact that antimanic treatment, in other words psychotrope use was not controlled for is an important limitation of the present study. Antimanic treatments received by our cases is as follows: lithium in 8 cases, sodium valproate in 12 cases, lithium and ketipin in 7 cases, lithium and olanzapine in 3 cases, and sodium valproate and olanzapine in 8 cases. Another limitation of our study was that there was no healthy control group.

Table 1. Sociodemographical and clinical properties of BD patients

<table>
<thead>
<tr>
<th>Gender (F/M)</th>
<th>Age (Mean±SD)</th>
<th>Education year (Mean±SD)</th>
<th>Age of onset (Mean±SD)</th>
<th>Family history (%)</th>
<th>Severity of episode (YMRS), (Mean±SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>22/28</td>
<td>22.6±3.7</td>
<td>7.8±1.1</td>
<td>22.6±3.7</td>
<td>46</td>
<td>28.3±3.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>32</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>46.7±11.5</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>55.6±20.9</td>
</tr>
</tbody>
</table>

Table 2. Comparison of peak power values between mania and remission

<table>
<thead>
<tr>
<th>Mania n=38</th>
<th>Remission n=38</th>
<th>Analysis p</th>
</tr>
</thead>
<tbody>
<tr>
<td>FP2-F4</td>
<td>0.1852±0.1195</td>
<td>0.0891±0.0452</td>
</tr>
<tr>
<td>F4-C4</td>
<td>0.0873±0.0724</td>
<td>0.0583±0.0161</td>
</tr>
<tr>
<td>C4-P4</td>
<td>0.0897±0.0703</td>
<td>0.0557±0.0306</td>
</tr>
<tr>
<td>P4-O2</td>
<td>0.0927±0.0841</td>
<td>0.0653±0.0299</td>
</tr>
<tr>
<td>FP2-F8</td>
<td>0.1613±0.1429</td>
<td>0.1712±0.1662</td>
</tr>
<tr>
<td>FB-T4</td>
<td>0.1663±0.1662</td>
<td>0.1603±0.1484</td>
</tr>
<tr>
<td>T4-T6</td>
<td>0.1432±0.1374</td>
<td>0.1435±0.1299</td>
</tr>
<tr>
<td>T6-O2</td>
<td>0.1439±0.1075</td>
<td>0.1390±0.1335</td>
</tr>
<tr>
<td>FP1-F3</td>
<td>0.1665±0.1424</td>
<td>0.1322±0.0866</td>
</tr>
<tr>
<td>F3-C3</td>
<td>0.1016±0.0883</td>
<td>0.0753±0.0703</td>
</tr>
<tr>
<td>C3-P3</td>
<td>0.1415±0.1247</td>
<td>0.1008±0.0907</td>
</tr>
<tr>
<td>P3-O1</td>
<td>0.1185±0.1085</td>
<td>0.1102±0.1065</td>
</tr>
<tr>
<td>FP1-F7</td>
<td>0.1430±0.1204</td>
<td>0.1119±0.0538</td>
</tr>
<tr>
<td>F7-T3</td>
<td>0.1350±0.1078</td>
<td>0.0687±0.0445</td>
</tr>
<tr>
<td>T3-T5</td>
<td>0.1588±0.1003</td>
<td>0.1359±0.1246</td>
</tr>
<tr>
<td>T5-O1</td>
<td>0.1373±0.0676</td>
<td>0.1336±0.1246</td>
</tr>
</tbody>
</table>

Tests with emotional content, beta activity increased, particularly in the temporal region. In the present study, in first-episode mania cases, unlike in the remission period, higher maximum peak values were obtained from right frontotemporal and left frontotemporal electrodes. Beta band corresponding to these values is responsible for maintenance of present sensorimotor and cognitive status (status quo) and for preparation of (go–no go) responses and inhibitor control (17). Pathological changes in the activity of this band are linked to abnormal resistance in present condition, and lack of flexible cognition and behavior (18). EEG spectral power analysis has revealed that significant effect was generally observed in beta subbands with low frequency and that the highest peak power values belonged to them in performance tests.

Findings supporting the above findings can be found in studies on ADHS. In ADHS, beta activity was found to be higher than that in healthy controls, particularly in the frontal region, and it was reported that it decreased with drugs but did not return to normal and was associated with hyperactivity and impulsivity (19). In ADHS, frontal beta activity is higher in cases responding to stimulants than in cases not responding (20). In addition, frontal beta activity was found to be associated with improvement in Conners Continuous Performance Test performance (maintenance of attention and inhibitor control) and decrease in hyperactivity–impulsivity scores reported by the family. Increased beta oscillatory beta activity after a positive feedback has been repeatedly demonstrated in previous studies (21). In gambling test including both positive and negative feedback, unexpected awards were markedly associated with increased beta activity (22).
Despite the limitations, in the present study, important electrophysiological data and clinical connections regarding BD were disclosed. First-episode mania and the subsequent remission period display differences electrophysiologically. These differences are characterized by abnormal resistance and lack of flexible cognition and behavior. In addition, peak power values of PSD during the remission period aid in distinguishing cases with psychotic findings in manic period from those without psychotic findings. Our cases were first-episode cases indicated that electrophysiological corollary of BD is present starting from the onset of disease.

Conflict of Interest: No conflict of interest was declared by the authors.

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