Late Positive Potential in Bipolar Disorder Reflecting Emotional Disorder: An Event-Related Potential Study

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ABSTRACT

Introduction: Bipolar disorder (BD) is an emotional disorder characterized by emotional instability. This study aims to investigate whether emotional disorders affect the cortical processing that regulates emotional stimuli.

Methods: This study uses a passive viewing paradigm to compare event-related potentials (ERPs) of 21 subjects with bipolar disorder (BD), 25 subjects with schizophrenia (SZ), and 25 normal subjects (HC).

Results: In the frontal and parietal areas (1) the late positive potential (LPP) amplitude of BD was greater than that of SZ and HC, and there was no difference in terms of peak latency. (2) In the occipital area, the amplitude and latency of P300 of SZ and BD were significantly larger than HC, but there was no difference between SZ and BD. (3) The behavioral results demonstrated that BDs’ valence rating on both positive and negative stimuli were significantly higher than those of HC and SZ when viewing face images, while BDs’ arousal scores for positive stimuli were significantly lower than those of HC and SZ. BDs’ arousal score for negative stimuli was significantly higher than those of HC and SZ. When viewing non-face pictures, BDs’ positive valence rating on negative stimuli and arousal scores for positive stimuli were significantly higher than those of HC and SZ.

Conclusion: Bipolar patients have complete early emotional recognition, and their emotional abnormalities exist in the late evaluation of emotional stimuli, mainly occurring in the frontal and parietal areas.

Keywords: Bipolar disorder; Late Positive Potential (LPP); Emotional process; Emotional evaluation

Abbreviations: LPP: Late Positive Potential; BD: Bipolar Disorder; SZ: Schizophrenia; HC: Health Control; ERP: Event-Related Potentials; VAS: Visual Analog Scales; DSM-IV: Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition; HAMD: The Hamilton Depression Rating Scale; CAPS: The Chinese Affective Picture System

INTRODUCTION

Bipolar disorder (BD) is an emotional disorder characterized by emotional instability [1,2], the main symptoms of which are emotional processing disorders [3,4]. While previous studies have found that patients with BD have mood-recognition disorder during remission, which appears as defective emotion recognition, other reports suggest that patients may have complete emotional perception [5]. Townsend’s study found that BD includes emotional and attention-biased impaired identification, which means that patients with BD are biased about both positive and negative emotions and BD patients with depression are particularly negative regarding negative emotions. A previous study of P300 in bipolar mania has confirmed that mania is a serious emotional disorder with impaired emotional processes [6]. Emotional processing consists of emotional responses, including early emotional assessments and post-reviews [7]. This study aims to investigate whether emotional disorders affect the cortical processing that regulates emotional stimuli. In other words, it explores the temporal progression and temporal-temporal relationship of late emotional processing by investigating the event-related potential (ERP) differences in...
emotional processes between different groups of subjects; namely, BD and normal. Another objective of this study is to explore the ERP differences in emotional processes simulated by human faces and non-face images. Three hypotheses are proposed:

- There is no difference in the average amplitude of P300 across the three groups - BD, SZ, and HC; BD triggers a larger amplitude of late positive potential (LPP) than the other two groups.
- LPP is more likely to be caused by face images than non-face images as the LPP amplitude of non-face images was larger than that of face images, and the latency was earlier.
- BD has higher valence and arousal on neutral stimuli than the other two groups.

RESEARCH METHODOLOGY

Participants
Two psychiatrists divided subjects into two groups of 26 people each according to their DSM-IV diagnosis, as shown in Table 1. All subjects are right-handed and have no neurological disease, persistent loss of consciousness, or major illness. The healthy subjects have no history of head injury, family history, no alcohol or drug dependence, and no disorders. Written informed consents were signed on the day of the experiment. Emotional symptoms were tested in the BD group: mania symptoms were assessed using the Young Mania Rating Scale and depression severity was assessed using the Hamilton Depression Rating Scale (HAMD). Seven subjects (five in the BD group, one in the SZ group, and one in the normal group) were excluded from offline analysis as they had significant selection biases in behavioral data.

Table 1: Participant demographics.

<table>
<thead>
<tr>
<th>Group</th>
<th>Sex</th>
<th>Age(years)</th>
<th>Edu(years)</th>
<th>BRPS</th>
<th>HAMD</th>
<th>YMRS</th>
</tr>
</thead>
<tbody>
<tr>
<td>BD</td>
<td>N=26(f=11)</td>
<td>41.0±15.2</td>
<td>9.1±0.2</td>
<td>25.4±9.1</td>
<td>3.0±3.8</td>
<td>12.4±7.6</td>
</tr>
<tr>
<td>SZ</td>
<td>N=26(f=13)</td>
<td>44.3±12.3</td>
<td>9.8±4.2</td>
<td>33.5±10.5</td>
<td>11.0±3.6</td>
<td>8.3±12.7</td>
</tr>
<tr>
<td>HC</td>
<td>N=26(f=13)</td>
<td>43.1±14.1</td>
<td>9.1±3.5</td>
<td>~</td>
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</tbody>
</table>

Standard deviation in parentheses; BD: Bipolar disorder; SZ: Schizophrenia; HC: Healthy control group

Stimuli and procedure
A Passive Viewing Paradigm was adopted as the experimental paradigm and presented in E-prime 2.0. Thirty face photos (angry/happy/neutral) and 30 emotional pictures (unpleasant/happy/neutral) were displayed and repeated once. A total of 360 trials were conducted. The experiment consisted of four blocks, and the ABBA design was used to exclude the order effect. After each block, the rest time was determined by the subject, with a maximum pause of 2 minutes. The experimental flow is shown in Figure 1. After the EEG test, subjects were asked to rate their levels of valence and arousal for the emotional stimuli. According to Visual analog scales (VAS), the subjects used the mouse to select the percentage level (leftmost 100, middle 50, rightmost 0) indicated on a horizontal bar. Prior to the experiment, 10 trials were conducted to confirm participants’ understanding of the process and the experiment then commenced [8].

The experiment was conducted in a comfortable room with hospital lighting and soundproofing. The distance between the screen and the eyes of the subject was 1m, and the stimuli were presented in the center of the screen.

Figure 1: Experimental design (A Passive Viewing Paradigm [12] was adopted as the experimental paradigm and presented in E-prime 2.0. Thirty face photos (angry/happy/neutral) and 30 emotional pictures (unpleasant/happy/neutral) were displayed and repeated once. A total of 360 trials were conducted. The experiment consisted of four blocks, and the ABBA design was used to exclude the order effect. After each block, the rest time was determined by the subject, with a maximum pause of 2 minutes.)

Face and non-face images were selected in the Chinese Affective Picture System (CAPS); the hairline of the face was removed, and the images were in black and white. The results of valence are: negative face (M=2.74, SD=0.47), negative picture (2.76, 0.62); positive face (6.56, 0.54), positive picture (6.70, 0.42); neutral face (4.72, 0.38), neutral picture (5.14, 0.42). The arousal is: negative face (6.50, 1.04), negative picture (5.75, 0.78); positive face (6.59, 0.63), positive picture (5.82, 0.34); neutral face (3.09, 0.31), neutral picture (3.88, 0.65).

ERP data recording and analysis
In the passive viewing paradigm, the EEG was acquired by a 64-channel EEGOTM EE-211 amplifier, a 64-channel Ag/AgCl electrode cap, and the EEGOTM system of the ANT Neuro BV (Enschede, The Netherlands). The electrode impedance was less than 5kΩ; the A/D Rate was 1000Hz; the reference electrodes were GND and CPz; eye electricity was not recorded. In the offline analysis, the average of M1, M2 was used as a reference point. The filtering standard was 0.1 ~ 30Hz, 24 slope [dB/oct]. Software-specific algorithms were used to correct ocular artifacts. The default artifact removal was ±100 μV, and the total number of trials of subjects exceeded 80%. A segment of data from -200 to 2000ms was selected. In accordance with previous studies [9-11], Fz, Cz, Pz, CP1, CP2, PO3, and PO4 were selected to
measure the average amplitude and latency of ERP for the study of LPP. The time window was 400 ~ 1000ms. For all analyses, a Greenhouse-Geisser correction was applied to p values where appropriate.

RESULTS

Behavioral data

The main effects between groups were significant in the rating of valence, where F (2,67) = 6.704, p<0.01, and ηp2 = 0.167. The main effect of stimulus types was significant, where F (1,67) = 51.071, p<0.01, and ηp2 = 0.433. The main effect of emotional valence was significant, where F (2,134) = 262.614, p<0.01, and ηp2 = 0.797. Stimulus types, emotional valence, and interaction between groups were significant, where F (4,134) = 2.614, p<0.05, and ηp2 = 0.072. The simple effect analysis found that when watching face images, the BD group rated higher scores for positive valence to positive stimuli (36.44±8.84) than the normal group (27.29±14.09) and the SZ group (27.97±10.35). In terms of negative stimuli, the BD group’s scores for positive valence (70.36±15.88) were significantly higher than the normal group (55.06±9.18) and the SZ group (60.26±8.99). With regard to viewing images, the BD group’s scores for positive valence to the negative stimuli (56.80±19.20) were significantly higher than the normal group (48.53±5.04) and SZ group (38.11±16.27). No other effects of interest were significant.

The main effects between groups were significant in arousal ratings, where F (2,67) = 3.741, p<0.05, and ηp2 = 0.100. The main effect of emotional valence was significant, where F (2,134) = 7.164, p<0.01, and ηp2 = 0.176. The simple effect analysis found that when watching face images, the arousal score of the positive stimulation was significantly lower in the BD group (38.13±9.05) than in the HC group (58.41±24.11) and the SZ group (51.76±20.74). The BD group’s arousal rating of negative stimulation was significantly higher than that of the normal group (21.51±8.93) and the SZ group (24.07±9.55). With regard to viewing images, the BD group’s arousal rating of positive stimulation (72.76±15.66) was significantly higher than that of the normal group (45.47±23.16) and the SZ group (48.56±19.38). No other effects of interest were significant.

ERP analysis

As seen from the total average map (Figure 2), significant P300 and LPP were induced in certain parts of the brain in both groups. Significant slow wave (LPP) peaks caused by different groups of subjects were significantly different between approximately 400 ms and approximately 1000 ms.

Repeated measures analysis of variance (ANOVA) demonstrated significant interaction between brain areas and groups, where F3,89 = 3.908, p<0.05, and ηp2 = 0.102. The simple effect analysis found that the P300 amplitudes of SZ and BD were significantly larger than that of HC in the occipital area, but there was no difference between SZ and BD (MSZ=0.15 uV, MBD=0.25 uV, MHC=0.58 uV). Regarding the P300 latency, the main effect was significant between groups, where F (2,69) = 5.438, p<0.01, and ηp2 = 0.136. The P300 latency of SZ and BD is longer than HC, but there is no difference between SZ and BD (MSZ=254.82ms, MBD=251.22ms, MHC=237.93ms). The simple effect analysis found that under positive stimulation, the P300 latency of SZ was longer than that of HC, but there was no difference between SZ and BD, or between BD and HC (MSZ=254.72ms, MBD=251.22ms, MHC=237.93ms); under neutral stimulation, SZ and BD had longer latency than HC, and there was no difference between SZ and BD (MSZ=251.99ms, MBD=261.60ms, MHC=237.29ms).

Regarding the LPP amplitude, the main effect was significant between groups, where F (2,68) = 4.301, p<0.05, and ηp2 = 0.112. The LPP amplitude of BD was larger than HC (IJ=0.547, SD=0.186, p<0.01, but did not differ greatly from SZ (IJ=0.316, SD=0.186, p>0.05), and there was no significant difference between SZ and HC (IJ= 0.229, SD=0.178, p>0.05). The main effect of brain region was significant, where F2,136 = 28.838, p<0.01, ηp2 = 0.298. The interaction between brain region and group was significant, where F (4,136) = 3.902, p<0.01, ηp2 = 0.103. The simple effect analysis found that in the frontal area, the LPP amplitude of BD was larger than that of HC and SZ, and there was no difference between SZ and HC (MSZ= -0.085 uV, MBD= 0.454 uV, MHC= -0.137 uV); in the parietal area: the LPP amplitude of BD was greater than HC and SZ, and SZ was significantly larger than HC (MSZ= -0.281 uV, MBD= 0.250 uV, MHC= -0.591 uV).

No other effects of interest were significant.
DISCUSSION

Previous studies on patients with bipolar disorder have found that bipolar patients had emotional processing disorders during emotional processes. Based on this theoretical hypothesis, this study explored the differences between people with bipolar disorder, schizophrenia, and normal people, and the ERP effects of different types of stimuli.

The most important finding of this study is that in the frontal and parietal areas, the mean amplitude of the LPP induced by BD patients was greater than that of normal and schizophrenic patients, and there was no difference in the mean amplitude of P300. Thus, our hypothesis is validated. The P300 volatility is related to the allocation of attentional resources [6,12,13]. The results indicate that the attentional resource allocation of the BD group was good and the emotional perception might be complete. However, related studies have found that BD patients with episodes had reduced P300 amplitude under negative stimulation. Since this contradicts our experimental results, further verification is required. In addition, in the valence rating, when viewing face images, the BD’s valence score was higher than those of the control groups. This result is consistent with the LPP volatility results but does not match the P300 volatility results. The results of this behavior are in line with the rating results of Degabriele [14], which found that the BD group had higher positive valence and arousal ratings on neutral stimuli. It should be noted, however, that compared with the normal group, the BD’s higher scores were not only on a single neutral stimulus but on all face images, which may be related to patients’ emotional state. Therefore, we believe that bipolar patients have a certain degree of emotional processing disorders, which are characterized by sustained attention and processing of emotional stimuli and increasing LPP amplitude at late stage.

This study found that all three groups demonstrated similar LPP amplitude, but there was a difference in terms of the frontal and parietal areas. The BD group induced a larger LPP amplitude, perhaps because the initial emotional perception involved the occipital lobe, and the late emotional processing brain areas are mainly subcortical or cortical-marginal systems. Moratti, Saugar, and Strange also ascended to the parietal and frontal cortex, and believed that to be the basis for the production and maintenance of LPP [14]. However, other studies have traced LPP production and maintenance back to the visual attention area in the back of the brain [15].

In addition, we found that with both positive and neutral stimuli, the BD’s P300 latency was later than the normal group, while there was no significant difference in the LPP latency between the BD and normal subjects. The study believes that the classified stimuli increased the latency of the P300, which is the time required for the P300 latency to depend on classified stimuli. Therefore, the prolongation of the BD’s P300 latency may be related to the greater attention paid by BD patients to the stimulation. There was no difference in LPP latency between BD and normal patients. The LPP demonstrated high and low sensitivity to the arousal of stimuli, indicating the similarity of the arousal between the BD and normal groups. However, the results of behavioral arousal ratings do not support this.

The comparison of the average amplitudes of P300 and LPP demonstrate that there was no difference in the P300 amplitude between groups but the LPP differed across groups. The LPP represents the persistence of late emotional processing of attention, meaning that LPP can better reflect emotional processing disorders than P300.

In this study, no hemispherical lateralization effect was observed in either the average amplitude or peak latency of the LPP. Previous studies [15] suggest that lateralization is related to emotional valence, where the left brain is more sensitive to positive stimuli and the right brain to negative stimuli. Other studies, such as a study of phobia [16], have also suggested that lateralization is generally associated with individual differences in gender, disease, etc. However, the lateralization effect has not been reported consistently, and this is still the case in the passive viewing paradigm, so the hemispheric effect should be reported cautiously. For example, one study by Horan et al. [12] reported that normal people had higher LPP in the left brain than in the right regarding the stimulation of valence, but the lateralization effect was not found for the LPP amplitude of schizophrenic patients. Similar findings have also been mentioned in Cunningham [17]. These results match the neuropsychological model of processing positive emotion with approach motivation, but whether it is approach or avoidance motivation for negative stimuli remains controversial [18,19]. For instance, recent studies have demonstrated that anger is an approach motivation with more stimulation in the frontal cortex in the left hemisphere than in the right hemisphere. However, the hemispheric effect was not found in the LPP in this study. The lateralization may be related to motivation. It may indicate that the stimulation was not sufficiently strong in the passive viewing paradigm and thus led to the subject’s strong approach or avoidance motivation. Further study of the hemisphere lateralization effect is still required.

Emotional stimuli are commonly used to trigger LPP. Many studies have demonstrated that emotional stimuli and neutral stimuli can induce different LPP. For example, positive and negative stimuli usually trigger larger LPP waves in a sequence task that presents positive-neutral-negative stimuli. In previous research, Caccioppo et al. found that emotional stimuli can increase the amplitude of LPP and can arouse the organism if the strength is sufficiently strong, as neutral images do not have this effect. Later studies have continued to confirm that positive and negative pictures as well as words can cause large LPP amplitudes compared with neutral pictures and words [19].

Unexpectedly, this study only observed that the P300 amplitude for negative stimuli was greater than that for positive and neutral stimuli, and there was no difference between positive and neutral in the frontal and parietal areas, while LPP amplitudes for negative, positive, and neutral stimuli were similar. The results of emotional valence of P300 may suggest that P300 is sensitive to emotional stimuli. In terms of LPP, some scholars believe that emotional valence does not have specificity. For example, Schupp [20] found that at the top of the hindbrain, LPP does not have specificity of emotional valence. This means that the LPP waves for positive and negative stimuli are both high, without any difference between them. Furthermore,
Cunningham [21] also mentioned the problem that emotional valence has no specificity. The lateral motive hypothesis mentioned above also suggests that the effect caused by emotion is not related to emotional valence, and our experimental results have validated this hypothesis.

This study has a number of limitations. First, it used different diagnostic criteria and different measurement standards for the stabilization phase and episodes, which might inevitably affect the screening of subjects. Moreover, this study only investigated the emotion paradigm of BD subjects during manic episodes (highly-emotional group) to explore the mechanism of late emotional processing. Future research can conduct a comparative study of multiple emotional states of BD patients, such as patients that are highly-emotional, depressed, and in remission [5]. Additionally, considering that the use of psychotropic drugs may lead to cognitive dysfunction, it is necessary to control or monitor the use of test drugs in the experiment in future studies, as Degabriele [14] mentioned that it was important to monitor the effects or influence of psychotropic drugs.

CONCLUSION

The following conclusions can be drawn from the above discussion:

(1) BD patients have emotional processing disorders, which are characterized by sustained attention and processing of emotional stimuli and increasing LPP amplitude of the frontal and parietal areas.

(2) The volatility of P300 does not differ from normal people, meaning that early emotional processing of BD patients is complete.

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AUTHOR CONTRIBUTIONS

All authors contributed substantially and according to the BMC Psychiatry guidelines to be recognized as authors. Study concept and design: Qiang Zhou. Data acquisition: Lijun Zhang. Identification and quality assessment of studies: Qiang Zhou and Jia Chen. Data analysis and interpretation: Jia Chen. Manuscript preparation: Qiang Zhou and Jia Chen. Manuscript editing: Qiang Zhou and Liang Ying. All authors have read and approved the final version of the manuscript.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

Participants provided written informed consent. The study was approved by Research Ethics Committee Review of Wenzhou Medical University (Reference number: 2017062).

CONSENT FOR PUBLICATION

Not applicable.

AVAILABILITY OF DATA AND MATERIALS

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

COMPETING INTERESTS

The authors declare that they have no competing interests.

REFERENCES


