



Case Control Study

Impaired implicit emotion regulation in patients with panic disorder: An event-related potential study on affect labeling

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Abstract

BACKGROUND

Panic disorder (PD) involves emotion dysregulation, but its underlying mechanisms remain poorly understood. Previous research suggests that implicit emotion regulation may play a central role in PD-related emotion dysregulation and symptom maintenance. However, there is a lack of studies exploring the neural mechanisms of implicit emotion regulation in PD using neurophysiological indicators.

AIM

To study the neural mechanisms of implicit emotion regulation in PD with event-related potentials (ERP).

METHODS

A total of 25 PD patients and 20 healthy controls (HC) underwent clinical evaluations. The study utilized a case-control design with random sampling, selecting participants for the case group from March to December 2018. Participants performed an affect labeling task, using affect labeling as the experimental condition and gender labeling as the control condition. ERP and behavioral data were recorded to compare the late positive potential (LPP) within and between the groups.

RESULTS

Both PD and HC groups showed longer reaction times and decreased accuracy

under the affect labeling. In the HC group, late LPP amplitudes exhibited a dynamic pattern of initial increase followed by decrease. Importantly, a significant group $\times$ condition interaction effect was observed. Simple effect analysis revealed a reduction in the differences of late LPP amplitudes between the affect labeling and gender labeling conditions in the PD group compared to the HC group. Furthermore, among PD patients under the affect labeling, the late LPP was negatively correlated with disease severity, symptom frequency, and intensity.

CONCLUSION

PD patients demonstrate abnormalities in implicit emotion regulation, hampering their ability to mobilize cognitive resources for downregulating negative emotions. The late LPP amplitude in response to affect labeling may serve as a potentially valuable clinical indicator of PD severity.

Key Words: Panic disorder; Implicit; Emotion regulation; Affect labeling; Late positive potential

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Core Tip: This study investigates neural mechanisms of implicit emotion regulation in panic disorder (PD) using event-related potentials. PD patients exhibit anomalies during an affect labeling task, including prolonged reaction times and reduced accuracy. Neurophysiological data indicate diminished differences in late positive potential (LPP) amplitude between affect labeling and gender labeling in PD, negatively correlating with disease severity, symptom frequency, and intensity. Resultantly, PD patients demonstrate impaired implicit emotion regulation, hindering cognitive resource mobilization for negative emotion downregulation. The late LPP amplitude in response to affect labeling may serve as a valuable clinical indicator of PD severity.

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INTRODUCTION

Panic disorder (PD) is a severe anxiety disorder characterized by spontaneous and recurring panic attacks and a constant fear of experiencing panic symptoms[1]. Epidemiological studies have reported a lifetime prevalence of PD at 4.7%[2]. Despite advancements in understanding PD, the neural mechanisms underlying its onset remain incompletely understood. Clinical observations and experimental evidence suggest that deficiencies in emotion regulation play a crucial role in the development of PD[3] because they contribute to heightened fear responses.

Emotion regulation involves individuals modifying their emotional responses in response to environmental demands, exerting control over the duration, intensity, and underlying characteristics of emotions[4]. Previous studies on emotion regulation have primarily focused on explicit strategies. In the context of PD, research on the neural mechanisms of emotion regulation has predominantly examined conscious strategies, revealing abnormal activation in the dorsolateral prefrontal cortex (DLPFC) and the ventromedial prefrontal cortex in individuals with PD during emotional regulation processes[3]. However, meta-analyses on repetitive transcranial magnetic stimulation (rTMS) targeting brain regions associated with conscious emotion regulation, such as the DLPFC, have not provided conclusive evidence regarding the efficacy of rTMS in treating PD[5,6].

According to Gross's model[7], emotion regulation strategies can be explicit (conscious) or implicit (unconscious). In clinical settings, it is observed that panic attacks in PD patients often occur suddenly and cease abruptly, with some patients even experiencing attacks during sleep[1,8]. There is no conscious emotion regulation process before or during panic attacks in patients. The catastrophic cognitive theory suggests that in PD, patients automatically evaluate both internal and external stimuli that are insufficient to elicit intense reactions as threat signals, leading to the initiation and maintenance of their anxiety[8]. Additionally, our previous findings indicate that PD patients exhibit anomalous mismatch negativity responses to acoustic and visual stimuli (both emotional and non-emotional), indicating abnormal automatic information processing in these individuals[9]. Therefore, we speculate that the core mechanism underlying emotion regulation dysfunction in PD patients may not originate from explicit abnormalities in emotion regulation but instead from implicit abnormalities in emotion regulation.

While the reported evidence suggests a significant role of implicit emotion dysregulation in the pathogenesis of PD[10], the specific mechanisms underlying these regulatory effects remain unclear. Affect labeling is an effective way to reduce unwanted emotions and the distress associated with negative events through putting feelings into words[11]. As the intent to reduce distress is not explicit, affect labeling has been conceptualized as a form of implicit emotion regulation that typically involves verbally labeling the emotional content of a facial stimulus, such as labeling an angry facial expression as "angry" or a fearful facial expression as "fearful"[11,12]. Although the regulatory effects of affect labeling on emotions have been established[11,13], the precise mechanism by which it reduces emotions, whether through

direct weakening or a dynamic process of initial enhancement followed by attenuation, remains unclear, especially in PD patients. Therefore, employing high-temporal-resolution event-related potentials (ERP) can help elucidate these questions.

The late positive potential (LPP) is a prominent component of ERP, typically occurring around 300 ms after stimulus onset. It is widely utilized in the study of emotion regulation[14], and its amplitude is known to increase with the subjective intensity of emotional experience, serving as an indicator of emotional regulation effectiveness. While previous LPP research suggests that affect labeling can reduce emotional experience[11,13], these findings are based on measuring the overall effect after labeling. Affect labeling tasks involve cognitive and motivation-related processing, and the LPP reflects changes in cognitive resources and motivation-related stimulus processing[15]. As a result, the amplitude of the LPP may dynamically vary instead of simply weakening.

To date, no studies have investigated the neural correlates of affect labeling in PD using ERP. In our current investigation, we recorded LPP data during affect labeling tasks performed by both PD patients and healthy individuals. We used emotional faces (negative and positive) as stimuli, building upon findings from research on implicit emotion dysregulation in anxiety disorders[16]. Building on clinical observations and previous studies, we developed two hypotheses: (1) In healthy individuals, the amplitude of the LPP is expected to dynamically fluctuate rather than simply decrease when using affect labeling to downregulate emotions; and (2) Compared to healthy controls (HC), individuals with PD are anticipated to show a deficit in the regulatory effect of affect labeling, potentially attributed to impaired cognitive processing.

MATERIALS AND METHODS

Participants

Twenty-five individuals (7 males, 18 females) diagnosed with clinically predominant PD according to the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV)[17], and 20 HC (9 males, 11 females) without a history of DSM-IV Axis I disorders were recruited from the outpatient and emergency departments of the First Affiliated Hospital of Dalian Medical University and the surrounding communities. The sample size was determined based on previously published ERP studies related to PD[9]. The PD group scored 14 or higher on the Hamilton Anxiety Rating Scale (HAMA) and 7 or lower on the Hamilton Depression Rating Scale (HAMD). Benzodiazepine medication was discontinued for at least one week, and a standardized Structured Clinical Interview for DSM-IV was conducted by a qualified physician to screen for other Axis I diagnoses. The HC group was recruited during the same period and matched the demographic characteristics of the PD group. The HC group scored 7 or lower on the HAMA and HAMD. PD patients completed the PD Severity Scale (PDSS)[18] and the Panic-Associated Symptom Scale (PASS)[19] to assess the severity of PD and associated symptoms. All participants scored ≥ 24 on the Mini-Mental State Examination and abstained from using psychoactive substances in the 24 h preceding the examination. Participants with other psychiatric disorders, severe physical illnesses, brain disorders, or substance dependence were excluded. Informed consent was obtained from all participants prior to their participation, following the principles of the Declaration of Helsinki. This study was approved by the Ethics Committee of the First Hospital of Dalian Medical University.

Stimulus materials

From the Chinese Facial Affective Picture System[20], a set of 60 negative emotional face pictures (including anger, fear, and sadness) and 60 positive emotional face pictures were selected. The distribution of male and female pictures was equal in both categories. Statistical analysis revealed no significant differences in valence and arousal ratings between male and female pictures ($t = -1.54$, $P > 0.05$; $t = 0.19$, $P > 0.05$). The mean valence ratings for positive male and female pictures were 5.6 (SD = 0.6) and 6.3 (SD = 0.7), respectively, while negative male and female pictures received valence ratings of 2.8 (SD = 0.3) and 3.0 (SD = 0.5), respectively. Arousal ratings showed that positive male and female pictures had mean scores of 4.6 (SD = 1.1) and 4.8 (SD = 1.0), respectively, whereas negative male and female pictures had mean scores of 6.1 (SD = 1.2) and 5.8 (SD = 1.2), respectively.

Experiment design and procedure

The experiment followed a classic paradigm[21] that involved two tasks: Affect labeling (experimental group) and gender labeling (control group). Participants first fixated on a "+" in the center of the screen for 1000 ms. Then, for 3500 ms, they viewed randomly presented label words and targeted emotional facial expressions. In the gender labeling task, participants determined the gender of the facial image and pressed the left or right key accordingly. The gender labels used were "male" and "female." The affect labeling task followed the same procedure, with participants selecting the label word that best described the emotion. Each trial followed the procedure depicted in Figure 1, with the next trial commencing after the image disappeared. The experiment comprised a practice phase and a formal experiment phase, with the practice images excluded from the formal phase. The practice phase continued until participants demonstrated comprehension of the experimental procedure and achieved an accuracy rate of 80% or higher. The formal experiment phase consisted of 240 trials, with 120 trials allocated to affect labeling (randomly divided into E1 and E2 groups) and 120 trials to gender labeling (randomly divided into S1 and S2 groups). Each set of trials constituted a block. To ensure balance between participant groups, the experiment was conducted in two versions: E1-S1-E2-S2 and S2-E2-S1-E1.

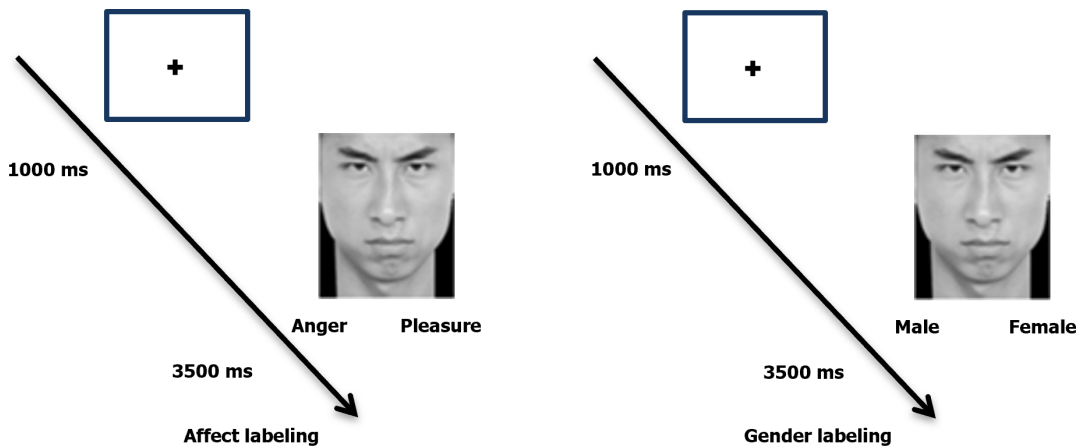


Figure 1 Task design. In the left panel, participants engage in affect labeling trials by selecting the suitable affect label that best describes the facial expression of the target face. In contrast, the right panel demonstrates gender labeling (control) trials, where participants are tasked with selecting the name corresponding to the gender-appropriate category of the target face.

Electroencephalogram recording

Stimulus presentation and behavioral response data (reaction time, accuracy) were controlled and recorded using E-prime 2.0 software. Electroencephalogram (EEG) data were collected using a Neuroscan SynAmps2 amplifier with a 64-channel Ag/AgCl electrode cap following the standardized 10-20 system. Continuous EEG signals were recorded and analyzed offline using NeuroScan 4.5 software. During EEG recording, an average reference was applied, and bipolar electrodes were used to capture horizontal eye movements and vertical eye movements. The data were sampled at a rate of 500 Hz in direct current mode and filtered with a bandpass of 0.05 to 100 Hz. Electrode impedance was maintained below 5 k Ω throughout the data collection process.

Data analysis

The participants' EEG waveforms were analyzed offline using NeuroScan 4.5 software. Bilateral mastoids served as the average reference electrodes. Manual removal of eye movements and artifacts was performed. The EEG data were filtered with a low-pass filter at 30 Hz and a high-pass filter at 0.5 Hz. Segmentation of the EEG data yielded two time windows: Early LPP and late LPP. The early LPP time window extended from 200 ms before stimulus onset to 1000 ms after stimulus onset, while the late LPP was defined as the time window spanning from 200 ms before response onset to 1000 ms after response onset. Artifacts were rejected, and the data were averaged to obtain the grand average waveform. The measured ERP components, based on prior literature, comprised the early LPP (300-800 ms after stimulus onset, reflecting emotional experience following stimulus presentation) and the late LPP (300-1000 ms after task response, reflecting emotional processing and experience during task performance). To investigate the dynamic changes of the late LPP, the late LPP was further divided into two time windows (350-630 ms, 630-1000 ms) based on the grand average waveform from this experiment. Five electrode sites along the midline of the frontal-parietal region were recorded: Frontal midline (Fz), frontocentral midline (FCz), central midline (Cz), centroparietal midline (CPz), and parietal midline (Pz).

The behavioral data (reaction time, accuracy) and EEG components were analyzed using SPSS 22.0. A 2×2 repeated measures analysis of variance (RMANOVA) was conducted on the behavioral data, with group (PD, HC) as the between-subjects factor and task type (affect labeling, gender labeling) as the within-subjects factor. The LPP amplitude data were analyzed using a $2 \times 2 \times 5$ RMANOVA, considering group, task type, and electrode location (Fz, FCz, Cz, CPz, Pz). Separate within-group RMANOVAs (2×5) were performed for the HC and PD groups, with task type and electrode location as the within-subjects' factors. Covariance analysis was applied, taking into account covariates based on the behavioral results. Effect sizes were measured with partial eta-squared (η_p^2). Greenhouse-Geisser correction was applied, when necessary, to adjust degrees of freedom and *P*-values for main effects and interaction effects. Pearson correlation analysis was used to examine the relationships between electrophysiological data, clinical data, and behavioral data in the PD group. Two-tailed tests were conducted, and statistical significance was set at $P < 0.05$.

RESULTS

Demographic and clinical data

Demographic and clinical characteristics of the HC and PD groups are presented in Table 1. There were no significant differences among the three groups in terms of gender, age, body mass index, and education. However, significant differences were observed in HAMA and HAMD scores ($P < 0.001$), with the PD group exhibiting the highest scores for both HAMA and HAMD.

Table 1 Demographics and clinical characteristics of the participants

	PD (<i>n</i> = 25)	HC (<i>n</i> = 20)	<i>t</i> / χ^2	<i>P</i> value
Age (yr)	47.20 ± 10.928	49.65 ± 12.766	0.681	0.500
Gender (male/female)	7/18	9/11	1.401	0.236
Education in years	13.52 ± 10.689	14.05 ± 12.037	0.156	0.877
BMI	24.89 ± 3.612	24.77 ± 3.222	0.120	0.905
HAMA	19.40 ± 4.830	2.95 ± 3.000	13.299	< 0.001 ^a
HAMD	12.76 ± 2.85	2.85 ± 1.599	10.581	< 0.001 ^a
PDSS	8.56 ± 3.906	N/A	N/A	N/A
PASS	6.84 ± 2.954	N/A	N/A	N/A

^a*P* < 0.001.

BMI: Body mass index; PD: Panic disorder; HC: Healthy controls; HAMA: Hamilton Anxiety Rating Scale; HAMD: Hamilton Depression Rating Scale; PDSS: Panic Disorder Severity Scale; PASS: Panic-Associated Symptom Scale; N/A: Not applicable.

Behavioral performance

Descriptive statistics for reaction time and accuracy of the two groups under the two conditions are presented in Table 2. Reaction time analysis showed no significant interaction between group and task type ($F_{1,41} = 1.209$, $P = 0.278$, $\eta_p^2 = 0.029$). Task type had a significant main effect ($F_{1,41} = 18.54$, $P < 0.001$, $\eta_p^2 = 0.311$), indicating longer reaction times in the affect labeling task. The main effect of group was not significant ($F_{1,41} = 2.339$, $P = 0.134$, $\eta_p^2 = 0.054$), despite the PD group having longer reaction times in both tasks.

Accuracy analysis showed no significant interaction between group and task type ($F_{1,41} = 0.453$, $P = 0.505$, $\eta_p^2 = 0.011$). The main effect of group was not statistically significant ($F_{1,41} = 0.468$, $P = 0.498$, $\eta_p^2 = 0.011$), indicating similar accuracy between the two groups. However, there was a significant difference between the two groups in the affect labeling and gender labeling conditions ($F_{1,41} = 19.174$, $P < 0.001$, $\eta_p^2 = 0.319$), with higher accuracy in the gender labeling task.

Early LPP amplitude

The average early LPP amplitudes for both groups under the two conditions are shown in Supplementary Table 1, with waveform plots in Figure 2A and scalp topography in Figure 2B. Covariance analysis, with accuracy and reaction time as covariates, showed no significant differences in early LPP amplitudes among the groups, paradigm types, electrode locations, and their interactions ($P > 0.05$ for all comparisons).

Late LPP amplitude

As depicted in Figure 3A, the HC group demonstrated higher amplitudes compared to the PD group at FCz, Cz, CPz, and Pz electrode sites. We conducted an analysis of the overall average late LPP amplitude (300-1000 ms). Within the HC group, the affect labeling condition exhibited larger LPP amplitudes than the gender labeling condition [(2.77 ± 2.75 μV) *vs* (2.15 ± 2.42 μV)]. Conversely, in the PD group, there was minimal difference in late LPP amplitudes between the affect labeling and gender labeling conditions [(1.49 ± 2.29 μV) *vs* (1.79 ± 2.63 μV)], which aligns with the observed scalp voltage distribution in Figure 3B. Supplementary Table 2 provides the statistical values for the overall late LPP amplitudes.

A 2 × 2 × 5 RMANOVA was conducted to analyze overall average late LPP amplitude within the 350-630 ms time window. The results revealed significant effects. Firstly, there was a significant main effect of electrode location ($F_{1,46,63} = 33.68$, $P < 0.001$, $\eta_p^2 = 0.4394$). Specifically, the FCz electrode showed significantly higher amplitudes compared to the Fz, Cz, CPz, and Pz electrodes ($P < 0.01$ for all comparisons). Furthermore, the analysis indicated significant differences in late LPP amplitudes between the two groups at the Cz ($P = 0.015$), CPz ($P = 0.01$), and Pz ($P = 0.004$) electrode sites.

Importantly, there was a significant task type × group interaction effect ($F_{1,41} = 6.32$, $P = 0.02$, $\eta_p^2 = 0.14$) within the 350-630 ms time window. Follow-up RMANOVA on the HC group revealed that affect labeling LPP (3.88 ± 3.60 μV) was significantly higher than gender labeling LPP (2.33 ± 3.48 μV) ($P = 0.044$). However, in the PD group, the difference between affect labeling LPP (3.14 ± 3.65 μV) and gender labeling LPP (2.44 ± 4.04 μV) did not reach statistical significance ($P > 0.05$), suggesting a lack of affect labeling effect. Late LPP amplitudes within the 630-1000 ms time window exhibited no significant differences in late LPP amplitudes across the groups, paradigm types, electrode locations, and their interactions ($P > 0.05$ for all comparisons).

Notably, in healthy individuals, the affect labeling LPP amplitudes were significantly higher than the gender labeling amplitudes within the 350-630 ms window, which does not indicate a decline in emotional response. However, after 630 ms, there was a trend towards convergence between the affect labeling and gender labeling LPP amplitudes. Further analysis in the 630-1000 ms window revealed a smaller difference in LPP amplitudes between affect labeling and gender labeling [(1.65 ± 2.35 μV) *vs* (1.36 ± 2.16 μV)], compared to the 350-630 ms window [(3.88 ± 3.60 μV) *vs* (2.33 ± 3.48 μV)], with no statistically significant difference of the 630-1000 ms window LPP amplitudes ($F_{1,19} = 1.06$, $P = 0.316$, $\eta_p^2 = 0.05$), indicating a decline in emotional response. This suggests that the unique nature of affect labeling led to an initial increase followed by a decrease in LPP amplitudes.

Table 2 Accuracy and reaction time (ms) of the two groups under the two tasks

	PD				HC			
	Affect labeling		Gender labeling		Affect labeling		Gender labeling	
	M	SD	M	SD	M	SD	M	SD
Reaction time	0.9271	4.62	0.9594	4.85	0.9145	13.94	0.9382	14.21
Accuracy	1389.32	221.43	1239.97	275.19	1247.50	261.76	1158.9	255.87

PD: Panic disorder; HC: Healthy controls; M: Mean.

Correlations

We performed Pearson correlation analyses to investigate the associations between late LPP amplitudes (350-630 ms) at 5 electrode sites during affect labeling and demographic data, behavioral measures, and clinical scale scores in the PD group. As shown in Figure 3C, we found significant negative correlations between late LPP amplitudes and PDSS scores at the Cz electrode site ($r = -0.437$, $P = 0.029$). Additionally, there were significant negative correlations between late LPP amplitudes and PASS scores at the same Cz electrode site ($r = -0.423$, $P = 0.035$).

DISCUSSION

This study investigated the implicit emotion regulation capacity in individuals with PD using ERP and affect labeling paradigms, while also examining the dynamic changes in affect labeling effects observed in the HC. Our findings revealed that both the PD and HC groups demonstrated prolonged reaction times and reduced accuracy in the affect labeling task compared to the gender labeling task, potentially indicating a greater demand for cognitive resources during affect labeling[22]. Additionally, the LPP amplitudes in the affect labeling condition displayed an initial increase followed by a subsequent decrease, suggesting that the process of emotion regulation through labeling is not immediate. Importantly, we observed an interaction effect between group and task condition on the late LPP. However, in the PD group, affect labeling did not result in significant neural modulation, indicating the ineffectiveness of this strategy in reducing negative emotions among PD patients and highlighting compromised implicit emotion regulation abilities in PD. Lastly, the LPP amplitudes in PD exhibited a negative correlation with PDSS and PASS scores, implying that the inability to effectively attenuate negative emotions during implicit emotion regulation may be associated with disease severity, episode frequency, and symptom intensity.

The behavioral data showed that, in the HC group, reaction times were longer for affect labeling compared to gender labeling, while accuracy was higher for gender labeling, which is in line with previous findings[23]. This performance difference can be attributed to task difficulty[24]. Moreover, the RMANOVA analysis demonstrated a significant main effect of condition but did not reveal a main effect of group, indicating that individuals with PD do not exhibit differences in reaction times and accuracy compared to HC.

The dynamic changes in LPP amplitudes during affect labeling, characterized by an initial increase followed by a decrease, reflect a distinct psychological process worthy of discussion. Torrisi *et al*[22] (2013) discovered that individuals utilize the prefrontal-amygdala circuit to suppress emotions during affect labeling, leading to the downregulation of negative emotional experiences[22]. This process involves the consumption of cognitive resources. Our study involved participants translating perceived facial emotions into verbal expressions through the affect labeling task, which required semantic processing of emotional features and the utilization of cognitive resources[23]. While both affect labeling and gender labeling tasks involve transforming features into language, affect labeling specifically deals with emotionally arousing information, representing a more complex cognitive processing[11] that demands higher cognitive resource consumption. Consequently, this is likely to lead to a more pronounced increase in LPP amplitudes during affect labeling. Furthermore, the decline in LPP amplitudes, reaching a level comparable to or even lower than gender labeling, indicates the regulatory influence of affect labeling on emotions[11,12]. However, the absence of a significant decrease in LPP amplitudes after affect labeling in this study can be attributed to the relatively lower level of arousal elicited by the negative images employed[23,25].

In this study, we observed an interaction effect between group and task type, indicating that the relationship between these factors influenced the outcomes. Specifically, when analyzing the late LPP amplitudes in the PD group, we found no significant differences between affect labeling and gender labeling tasks. These findings suggest a lack of effect of affect labeling on emotion regulation in individuals with PD. Previous research has consistently reported cognitive biases and interpretive tendencies in individuals with PD[8,26], where they exhibit a tendency to prioritize attention towards threatening stimuli and engage in catastrophic interpretations of emotional cues. This cognitive bias hinders effective downregulation of negative emotions, as individuals with PD tend to persevere on negative aspects and struggle to mobilize sufficient cognitive resources for successful regulation of negative emotions. Furthermore, prior functional magnetic resonance imaging (fMRI) studies have consistently demonstrated compromised top-down unconscious cognitive control from the prefrontal cortex to the amygdala in individuals with PD[3,27]. Our study aligns with these findings as we did not observe a significant enhancement in the late LPP amplitude (350-630 ms) in the electro-

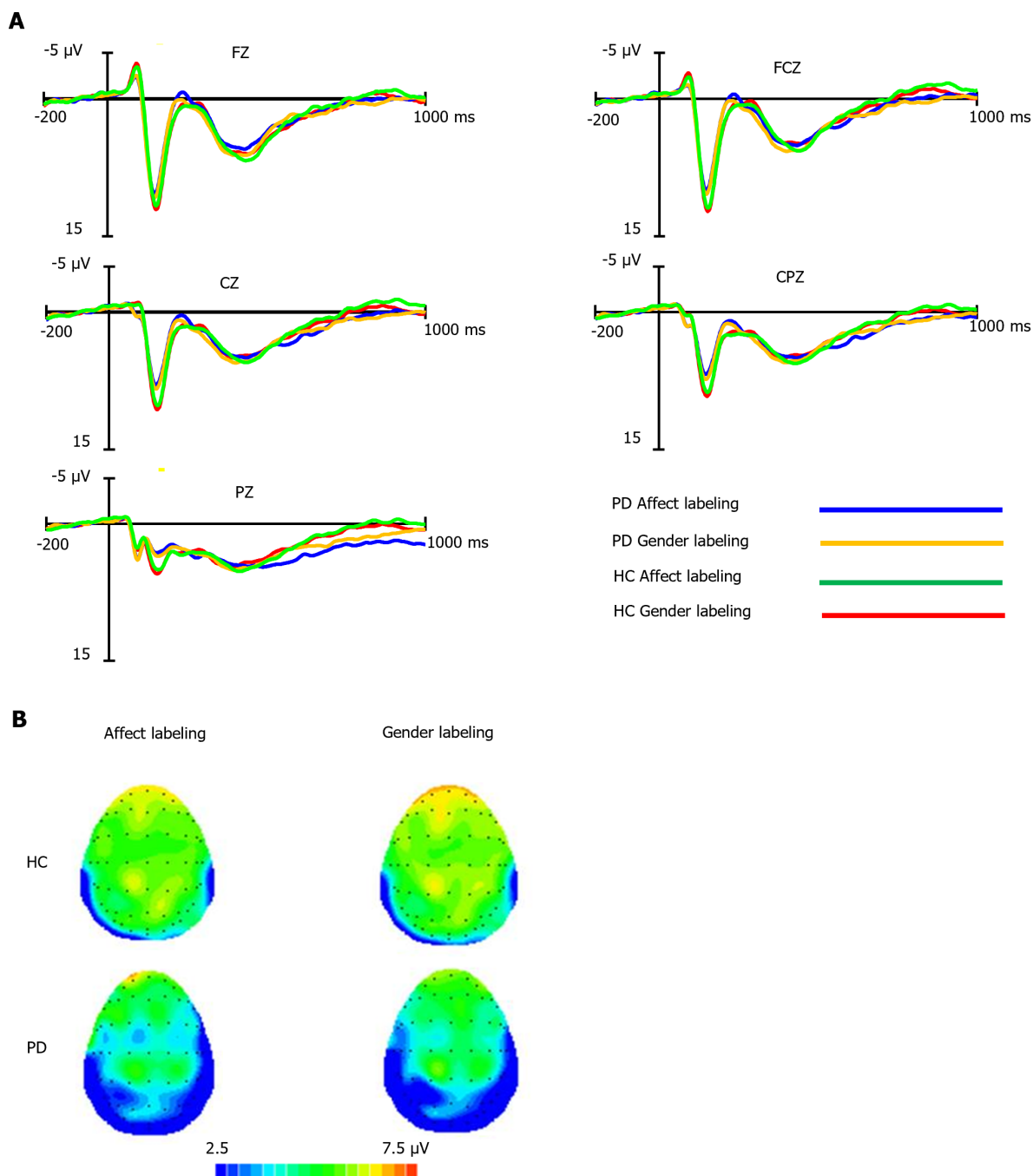


Figure 2 Early late positive potential amplitude. A: Illustrates the early late positive potential amplitudes of each electrode site for the healthy control and panic disorder groups under the affect labeling and gender labeling conditions. There were no significant main effects of group, condition, or electrode, and no significant interaction effects were observed ($P > 0.05$ for all comparisons); B: Displays the topographic maps of the two groups under the two conditions. PD: Panic disorder; HC: Healthy control; Fz: Frontal midline; FCz: Frontocentral midline; Cz: Central midline; CPz: Centroparietal midline; Pz: Parietal midline.

physiological signals, providing further support for the notion of impaired implicit emotion processing in individuals with PD.

Interestingly, we observed a significant negative correlation between disease severity and late LPP amplitudes in the PD group. Specifically, greater severity of PD symptoms in the past month or higher frequency and intensity of symptoms in the past week were associated with impaired implicit emotion regulation abilities. These findings support the potential use of LPP as an electrophysiological marker for monitoring disease progression and assessing treatment effectiveness.

This study has a few limitations. Firstly, the level of arousal elicited by the emotional stimuli was not strong enough, potentially impacting the full capture of the decline in late LPP[23,24]. However, data and waveform analysis support the existence of the regulatory process involving enhancement followed by attenuation through affect labeling. Secondly, the study exclusively utilized "negative" labels. Future research can explore more specific labels (e.g., neutral and positive) to further investigate related aspects. Lastly, the study design lacked longitudinal treatment evaluations. Future studies should consider incorporating follow-up assessments of the patients to track changes over time. This would not only

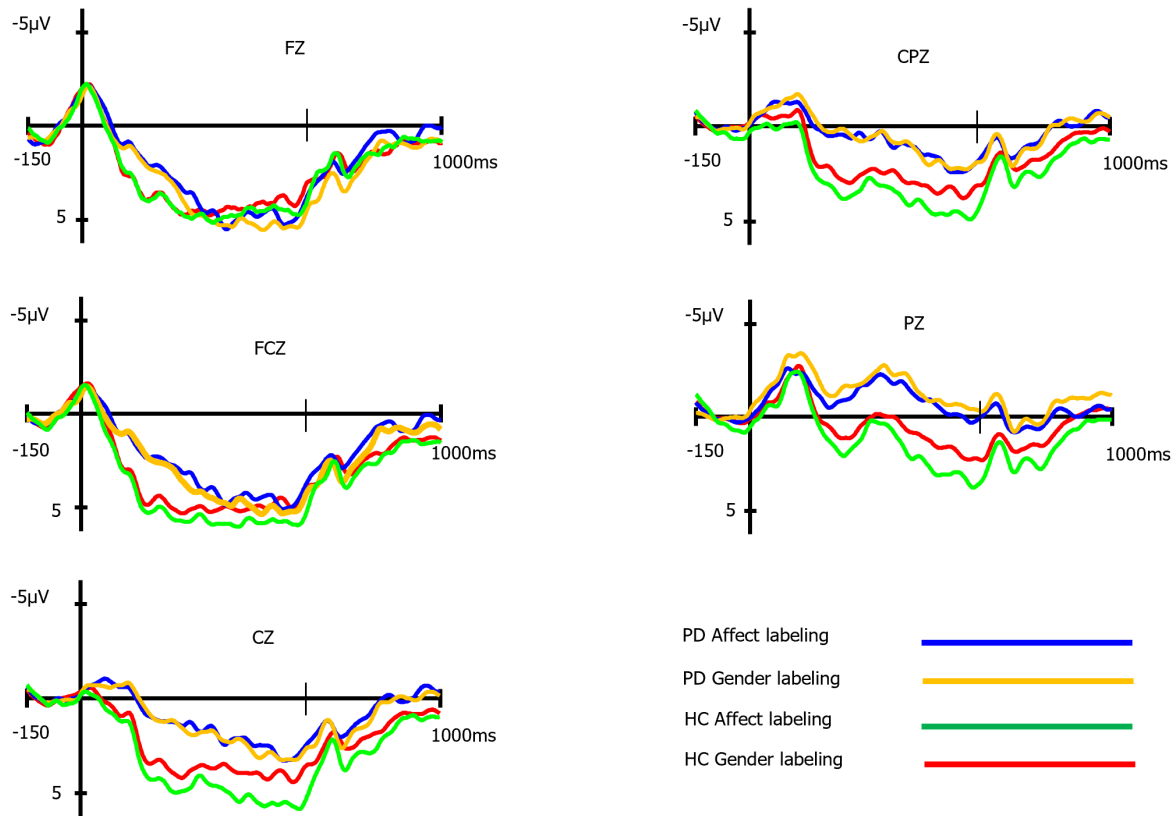
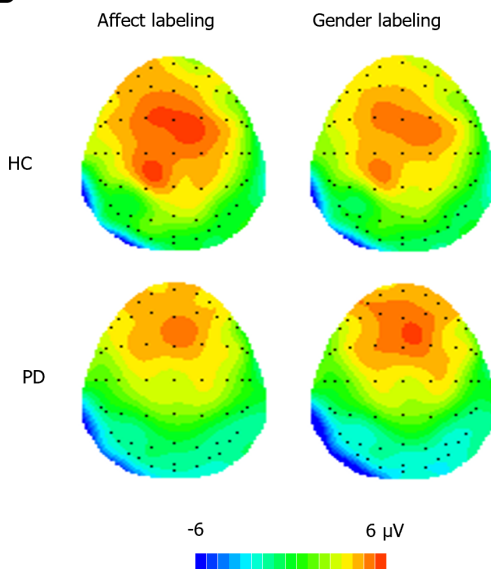
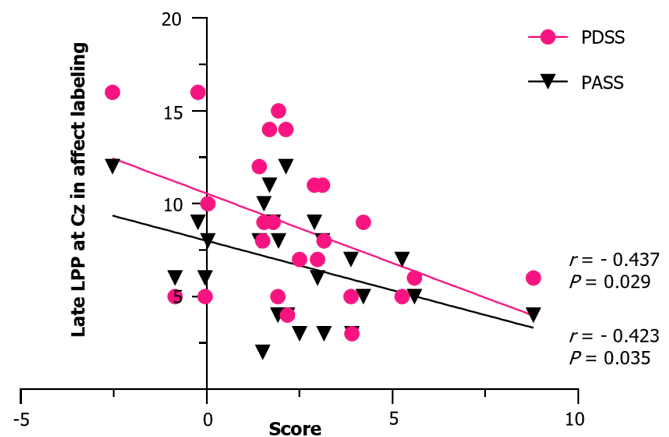
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Figure 3 Late positive potential amplitude. A: Illustrates the early late positive potential (LPP) amplitudes at each electrode site for the healthy controls (HC) and panic disorder (PD) groups under affect labeling and gender labeling conditions. Firstly, a $2 \times 2 \times 5$ repeated measures analysis of variance (350-630 ms) revealed significant main effects of electrode ($P < 0.001$) and a significant group $\times$ condition interaction effect ($P = 0.02$). Post-hoc comparisons indicated that the HC group exhibited higher LPP amplitudes in the affect labeling condition compared to the gender labeling condition ($P = 0.044$), while there was no significant difference in the PD group ($P > 0.05$). Secondly, no significant effects of electrode, condition, group, or interactions were observed in the 630-1000 ms time window ($P > 0.05$ for all comparisons); B: Presents topographic maps of the two groups under the two conditions; C: Demonstrates negative correlations between LPP amplitudes at the Cz site during affect labeling and Panic Disorder Severity Scale scores as well as Panic-Associated Symptom Scale scores in PD patients ($P = 0.029$, $P = 0.035$, respectively). PD: Panic disorder; HC: Healthy control; PDSS: Panic Disorder Severity Scale; PASS: Panic-Associated Symptom Scale; Fz: Frontal midline; FCz: Frontocentral midline; Cz: Central midline; CPz: Centroparietal midline; Pz: Parietal midline.

enhance our understanding of the causal relationship between affect labeling and neural activity changes in PD but also allow for an evaluation of whether successful treatment leads to the disappearance of the observed dysregulation.

CONCLUSION

This study reveals that the regulation process of affect labeling is not immediate, as evidenced by an initial increase followed by a decrease in LPP amplitude. Importantly, it identifies impaired implicit emotion regulation in PD patients and provides the first electrophysiological evidence of this impairment. Specifically, the diminished LPP modulation supports the abnormal affect labeling in PD patients. These findings serve as a valuable marker for future research investigating therapeutic interventions for PD that rely on implicit emotion regulation, such as cognitive behavioral therapy[28, 29], and neuromodulatory interventions like transcranial magnetic stimulation[6].

ARTICLE HIGHLIGHTS

Research background

The background of this study is rooted in clinical observations and research findings that identify emotion regulation dysfunction as a significant factor in the occurrence of panic disorder (PD). However, the neurophysiological mechanisms underlying implicit emotion regulation abnormalities in patients with PD remain unclear.

Research motivation

We aim to analyze neurophysiological changes in PD patients during implicit emotion regulation, identifying a concise and effective electrophysiological marker to assess potential anomalies in implicit emotion regulation.

Research objectives

The study aims to determine if there are anomalies in implicit emotion regulation in PD. Past research suggests abnormal Late Positive Potential (LPP) during emotion regulation in PD patients, indicating that LPP in event-related potentials (ERP) could be an effective tool for assessing implicit emotion regulation deficits.

Research methods

We assessed PD patients using clinical and psychological scales, conducting an emotion labeling task and recording behavioral and ERP data.

Research results

In the control group, late LPP initially increased, then decreased. A significant group $\times$ condition interaction effect was observed. Simple effect analysis showed reduced differences in late LPP amplitudes between affect labeling and gender labeling conditions in PD compared to controls. Additionally, under affect labeling, late LPP in PD negatively correlated with disease severity.

Research conclusions

PD patients have implicit emotion regulation impairments, and the late LPP amplitude in response to affect labeling may be a valuable clinical indicator of PD severity.

Research perspectives

Future considerations should include longitudinal treatment studies with follow-ups, assessing whether PD patients regain regulatory function post-successful treatment.

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FOOTNOTES

Co-first authors: Hai-Yang Wang and Li-Zhu Li.

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