INTRODUCTION

Electroencephalography (EEG) allows the non-invasive assessment of cognitive processes in real time [1]. The superior time resolution of EEG can follow fast neural events and permit the knowledge of related brain regions through source analysis [1]. It can directly assess the dynamics of neural activation related to various cognitive processes [2]. Moreover, it can be assessed in infants and children [3]; therefore, EEG was the first tool used to assess neural activities in children [4], and event-related potentials (ERPs) have been used in various studies to investigate psychopathology and neural substrates of psychiatric diseases in children [1,4].

This review aims to explain the current body of ERP studies on psychiatric illnesses in children and adolescents. It is important to understand how ERP studies have played a role in enhancing our understanding of neural functions in childhood and adolescence. This review starts with an introduction to the use of EEG and ERP to assess child-adolescent psychiatric diseases. Second, the definition of ERP and various ERP markers for child-adolescent psychiatric diseases will be outlined. Finally, recent advances in ERP research on representative child-adolescent psychiatric diseases will be reviewed.

Definitions of ERP

ERPs are scalp-recorded voltage changes that measure the response of the brain to a time-locked event [5]. In other words, ERPs allow us to determine the activities of coordinated neurons assessed by averaging sections of electrical recordings that are time-locked to motor, perceptual, and cognitive events [6]. The timing, size, and location of ERPs on the scalp infer the time course of neural processing in the brain [6]. Therefore, ERPs reflect thousands of simultaneously ongoing brain processes that can be used to evaluate perceptual processing, sensory-motor coupling, and cognition, such as attentional selection [7].

Use of EEG and ERP to assess child-adolescent psychiatric disease

ERPs enable a temporally detailed exploration of the cognitive process; early components of ERP reflect sensory or perceptual processing, and later components reflect higher
cognitive processes [8]. In particular, cognitive ERPs provide a powerful method for investigating synaptic functions in the brain. ERPs reflect excitatory postsynaptic potentials (EPSPs) and inhibitory postsynaptic potentials (IPSPs) [9-11]. Because ERP allows us to know the temporal patterns of neuronal activities, it can be useful in quantifying the sequence of various cognitive processes [12]. Considering that children and adolescents are in the phase of development, research involving ERPs will allow us to understand the association between brain development and behavioral development, which might provide important evidence [3].

There are many applications of EEG and ERP in child-adolescent psychopathologies within the context of biomarkers of disease [4]. Child-adolescent psychiatric illnesses are related to various factors, such as developmental course, clinically heterogeneous etiology, clinical manifestation, and treatment response. Therefore, the identification of biomarkers to facilitate diagnosis and predict treatment response is critical in this field. ERP could be a candidate biomarker in child-adolescent psychiatry, considering its ability to reflect cognitive and behavioral functions in humans.

From a practical perspective, it is crucial to use simple paradigms, especially for children. The stimuli for the ERP were relatively simple, such as a three-tone (standard, deviant, novel) auditory oddball paradigm [13]. Moreover, EEG and ERPs are non-invasive and inexpensive assessment methods; thus, they are useful for both clinical evaluation and research involving adolescents or children. Furthermore, the ERP is less affected by movement artifacts than other brain imaging methods, and it could be useful for studying children.

**Various ERP markers for child-adolescent psychiatric diseases**

**MMN**

Mismatch negativity (MMN) is an ERP component when a sequence of relatively standard stimuli is interrupted by the infrequent presentation of deviant stimuli [14]. In addition, MMN reflects preattentive auditory processing, which is closely correlated with cognitive status [15,16]. Regarding patients with psychiatric disease experiencing cognitive dysfunction, MMN has been considered a good tool for evaluating the neurophysiological mechanisms of psychiatric illness. Moreover, MMN reflects glutamatergic function [17], and the critical pathology of MMN reduction might be related to dysfunction of the N-methyl-D-aspartate (NMDA) receptor system [18]. NMDA receptors have been studied for their essential role in cognitive functions, such as learning and memory [19]. In this regard, MMN is considered a promising biomarker for many psychiatric disorders.

**P300, Go/NoGo paradigm, and Flanker task**

The P300 (P3) wave has two components, P3a and P3b. The P3a is a positive wave that appears maximally in the fronto-central region and has a peak latency of 250–280 ms [20]. It reflects the brain electrical activity related to decision-making, attention, and novelty processing [20]. The P3b component has a positive amplitude that peaks at approximately 300 ms; the peak will vary in latency from 250 ms to 500 ms or more. P3 was evoked using the oddball paradigm in which the target stimulus was mixed with a non-target stimulus. In particular, P3 can be elicited using various tasks, such as Go/NoGo and Flanker tasks. The Go/Nogo paradigm reflects behavioral and response inhibition [21,22]. NogoP3 is a well-known indicator of behavioral control [23]. Specifically, P3 reflects motor inhibition [24].

**ERN**

Error-related negativity (ERN) is a sharp negative amplitude that begins around the same time as an incorrect motor response [25], which occurs when a person makes an error [26]. It is evoked using cognitive tasks when a person has an incorrect response [27], and it appears as a negative wave at 80–150 ms following error commission [27]. Pe is a positive wave that appears subsequent to ERN and is the highest at 300 ms post incorrect response [26]. Although both Pe and ERN are associated with the error-related response, they are different in that Pe is related to post-error possessing [28], and ERN reflects the comparison process between an actual response and a required response [29]. That is, ERN can arise even after unaware errors, while Pe reflects conscious error processing [29]. Because ERN and Pe have different aspects of error processing, the two components have different neural generators [29]. Source localization indicates that ERN has a dipole in the anterior cingulate cortex, and Pe has a dipole in the posterior cingulate cortex [30].

**RewP**

Reward positivity (RewP) reflects reward responsiveness [31]. RewP is a positive wave following positive information and is either decreased or absent when obtaining negative information [32,33]. In other words, sensitivity to reward or non-reward stimuli can be represented by RewP [34]. RewP is the highest in medial fronto-central areas and appears at 250–550 ms post reward [31]. The amplitude of the RewP also predicts individual differences in sensitivity to reward, as assessed by both self-report and behavioral measures [35,36].
ERP markers for child-adolescent psychiatric disease

ADHD

Previous studies have discussed P3 latency in children with attention-deficit/hyperactivity disorder (ADHD). Participants with ADHD had an increased visual P3 latency and decreased P3 amplitude compared to healthy participants, suggesting reduced involvement in post-decisional processing [37]. Latency reflects attention, regarding it is various to the effort of discriminating different stimuli [38]. Latency is also associated with stimulus evaluation and discrimination [39]. Moreover, children with ADHD show increased latency and stimulants (methylphenidate) have been found to reduce the latency [38]. A previous study pointed out that children with ADHD have prolonged P3 latency at the fronto-central electrodes, and the level of prolongation also has a positive correlation with the severity of inattention [40]. This suggests that P3 latency is a neurophysiological marker in ADHD and may change under stimulant treatment [38].

Executive dysfunction is one of the well-known etiologies of ADHD, resulting in reduced cognitive function and abnormal behavior [41,42]. One of the most prominent etiological theories explains executive dysfunction as a secondary clinical manifestation of deficits in behavioral inhibition [43,44]. Specifically, the role of inhibition in ADHD has been emphasized by Nigg; the author insisted that ADHD is associated with deficits in behavioral inhibition [44-46]. On the other hand, error monitoring can show the self-monitoring of a person's own behavior, and it can make people assess their own response considering the actual demand. Therefore, it can recognize a potential error and help people adjust their response to prevent additional errors [44-46]. Patients with ADHD make many careless mistakes, and it could be expected that cognitive dysfunctions underlying error monitoring in various situations that require highly intended and about-to-be-performed responses may exist [44,45]. In this regard, error monitoring can be assessed using various tasks such as the Go/NoGo paradigm, stop-signal task, and Flanker task.

For the Go/NoGo paradigm, several previous studies reported that juvenile patients with ADHD have significantly different ERN compared to the healthy population [47-49]. In studies that used the Flanker task, both the Pe and Ne/ERN were decreased in patients with ADHD compared with the healthy population, and this finding was pronounced in the younger population [50].

One longitudinal EEG study that explored the course of impaired cognitive functions in ADHD has shown interesting results; all ERP components such as Cue P3, NoGo P3, and contingent negative variation (CNV) developed without significant time×group interactions [51]. Only CNV remained decreased in the ADHD group from childhood to adulthood [51]. This suggests that attentional and preparatory deficits in ADHD continue into adulthood, and attenuated CNV reflects a particularly stable ADHD marker. Further longitudinal studies are required to confirm these results [51].

Anxiety disorder

Anxiety disorder is one of the most common forms of child-adolescent psychopathologies and often begins early in human development [52]. ERN has been studied extensively in patients with anxiety disorders during childhood and adolescence. The ERN was shown to be increased in children with anxiety using various experimental paradigms to assess ERN [6,52,53]. Similar results were obtained using Flanker [6,52,53] and Go/NoGo tasks [54]. Ladouceur et al. [6] reported that children with anxiety disorder had increased ERN. In addition, a large ERN early in life (childhood and adolescence) was associated with an increased risk of developing anxiety disorder later in life [6,53]. Previous studies found that a larger ERN predicted the onset of anxiety disorder, while controlling for baseline anxiety symptoms and maternal history of anxiety [6,53,55].

A previous study that evaluated behavioral performance using the Go/NoGo task in a sample of 139 children aged 5–8 years reported that symptoms of separation anxiety disorder (SAD) were related to a small ERN, even after controlling for other anxiety disorder symptoms [56]. Additionally, children with more SAD symptoms showed higher error rates and failed to exhibit the expected association between behavioral performance and ERN, suggesting the presence of ineffective error-monitoring in children with SAD [56].

A recent prospective study showed meaningful results for the development of generalized anxiety disorder (GAD) [57] in 457 girls aged between 13.5 years and 15.5 years, with no history of GAD. The study revealed that the ERN using the Flanker task was a significant predictor independent of other risk factors, such as baseline anxiety and depression symptoms, and parental lifetime psychiatric history. This suggests the utility of ERN as a biomarker of GAD risk during a key developmental period.

Additionally, Carrasco et al. [53] revealed that the ERN magnitude was associated with obsessive-compulsive disorder (OCD) symptoms in children and adolescents without clinical OCD. This study reported that ERN was increased in unaffected siblings of children with OCD [53]. Moreover, a recent study found an increased ERN in children after successful treatment for OCD [58]. These studies suggest that children at risk for OCD may be characterized by increased ERN.
Mood disorders
Decreased motivation and reactivity toward pleasurable stimuli are the usual clinical manifestations of depression [59]. Depression onset often occurs in adolescence, the peak period for neural development of reward circuitry [42,60]. Blunted neural responses to RewP are related to vulnerability to depression [34]. A previous study with a community sample of 369 children showed that stressful life events moderated the effect of RewP on depression symptoms at follow-up [33]. In other words, blunted RewP predicted high risk of depression symptoms in individuals with high levels of stressful life events [33]. Another study with 20 adolescents aged 12–17 years reported direct effects of the RewP latency on depression symptoms and withdrawal behaviors [34]. In addition, longer reward latency is associated with stronger depression [34]. This suggests that avoidance motivation can help demonstrate the association between reduced reward responsivity and symptoms of depression [34].

CONCLUSION
Investigating ERPs is a noninvasive method to assess spatio-temporal activation in the brain during sensory, cognitive, affective, attentional, and motor information processing as well as during state regulation in real time. They can reliably be reproduced in a wide range of patients with psychiatric disease and across a wide age range, even in infants and children, and during both wakefulness and sleep. ERPs can be an ideal tool for studies of brain function during normal and deviant child development. Although more evidence related to various psychiatric conditions is needed, ERPs could be a candidate biomarker for studying child-adolescent psychiatry, considering their ability to reflect cognitive and behavioral functions in humans.

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Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

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