Modulation of Electrophysiology by Transcranial Direct Current Stimulation in Psychiatric Disorders: A Systematic Review

Minah Kim¹, Yoo Bin Kwak², Tae Young Lee³, and Jun Soo Kwon¹,²,³

¹Department of Psychiatry, Seoul National University College of Medicine, Seoul, Republic of Korea
²Department of Brain and Cognitive Sciences, Seoul National University College of Natural Science, Seoul, Republic of Korea
³Department of Neuropsychiatry, Seoul National University Hospital, Seoul, Republic of Korea

Objective Transcranial direct current stimulation (tDCS) is a non-invasive neuromodulation technique increasingly used to relieve symptoms of psychiatric disorders. Electrophysiologic markers, such as electroencephalography (EEG) and event-related potentials (ERP), have high temporal resolution sensitive to detect plastic changes of the brain associated with symptomatic improvement following tDCS application.

Methods We performed systematic review to identify electrophysiological markers that reflect tDCS effects on plastic brain changes in psychiatric disorders. A total of 638 studies were identified by searching PubMed, Embase, psychINFPO. Of these, 21 full-text articles were assessed eligible and included in the review.

Results Although the reviewed studies were heterogeneous in their choices of tDCS protocols, targeted electrophysiological markers, and disease entities, their results strongly support EEG/ERPs to sensitively reflect plastic brain changes and the associated symptomatic improvement following tDCS.

Conclusion EEG/ERPs may serve a potent tool in revealing the mechanisms underlying psychiatric symptoms, as well as in localizing the brain area targeted for stimulation. Future studies in each disease entities employing consistent tDCS protocols and electrophysiologic markers would be necessary in order to substantiate and further elaborate the findings of studies included in the present systematic review.

Key Words Brain plasticity, Electrophysiology, Neuromodulation, Psychiatric disorders, Transcranial direct current stimulation.
strate strengths. As measures directly reflecting neural activity during both resting and information processing states with high temporal resolution at the millisecond level, EEG and ERPs have long been employed in examining the altered neural processing associated with psychiatric disorders. These measures are also powerful tools for measuring plastic changes of brain after specific intervention, such as following medication, cognitive exercise, and tDCS. Therefore, investigation of tDCS effect on EEG and ERPs may provide further information on pathophysiologic mechanism underlying psychiatric disorders, as well as a serve as a guide for personalized neuromodulation in response prediction and selection of target areas. We performed the present systematic review to identify electrophysiological markers reflecting tDCS effects on plastic brain changes in psychiatric disorders, that, in turn, may serve as a biomarker for pathophysiology of psychiatric disorders.

METHODS

A systematic search strategy was used to identify relevant studies. Two independent researchers (M.K. and Y.B.K.) each conducted literature searches in PubMed, Embase, and PsycINFO databases for articles published during the period spanning from database inception to February 2017. The following Medical Subject Heading (MeSH) categories were used for the search: (electroencephalography OR EEG OR event related potential OR event-related potential OR ERP) AND (transcranial direct current stimulation OR tDCS). A total of 638 articles were found, and of these, only the studies satisfying the following criteria were included in the current review: 1) articles written in English; 2) original articles or short communications in peer-reviewed journals; 3) studies that included psychiatric patients; and 4) studies with outcome measured with EEG or ERP. The process for literature search and screening are presented in Figure 1.

RESULTS

Number of selected studies was 5 for schizophrenia, 5 for affective disorders, 6 for substance use disorders, 2 for Alzheimer’s disease (AD), and 3 for developmental disorders. Summary of studies pooled for review are shown in Table 1.

Schizophrenia

The 5 studies examining patients with schizophrenia applied anodal tDCS to modulate frontal cortical functions, although tDCS protocols used in each study showed slight variations. Of these, two studies examined tDCS effects on early and late auditory processing and reported inconsistent results. Alterations in the N1 and mismatch negativity (MMN), which reflect pre-attentive auditory processing, and in P3b or P300, which represents working memory update of change and attention, have all been reported in patients with schizophrenia. Knetchel et al. applied anodal tDCS over the left prefrontal cortex in 14 schizophrenia patients and examined effects on N1, MMN, and P3b ERP components. In this randomized cross over study, a single application of tDCS was found to have no acute effects on the ERP measures and associated auditory information processing in patients with schizophrenia. In a more recent randomized controlled study, Dunn et al. applied bilateral prefrontal tDCS on regions associated with auditory processing system in 36 patients with schizophrenia (12 patients in anodal, cathodal, sham group each) and examined effects on MMN and P300 ERP components. Compared to baseline, i.e., before stimulation, MMN amplitudes were found decreased post-stimulus for anodal tDCS. For P300 amplitudes, neither the main effects of time and condition nor their interaction were found significant. Results from Dunn et al. study suggest that bilateral prefrontal tDCS can engage neural systems underlying the impaired auditory processing in schizophrenia patients via the modulation of MMN, whose association with the pathophysiologic mechanism, clinical symptoms, as well as general functional status of schizophrenia are well known. Thus, although the two aforementioned studies did not measure or provide changes in psychotic symptoms following tDCS application, the reported MMN modulations following a single session of tDCS suggest tDCS as an potentially effective treatment option for improving symptoms and functioning in patients with schizophrenia.

The remaining three studies examined the effects of tDCS on higher cognitive functions, such as working memory, adaptive control, and learning ability, in relationship with modu-
Table 1. Summary of studies pooled for review

| Author, year | Study design | Patient number | tDCS protocol | EEG/ERP measurements | Conclusions |
|--------------|--------------|----------------|---------------|----------------------|-------------|
| Schizophrenia |              |                |               |                      |             |
| Knechtel et al., 2014 | Randomized cross over study | 14 SZ | Anode: left-prefrontal at F3 Cathode: right supraorbital region Stimilation: 2 mA, 20 min, single session "sham" and "active" tDCS took place 1 hour apart | N1, MMN, P3b (P300) | No effect of tDCS on any ERP component was detected |
| Dunn et al., 2016 | Randomized controlled study | 36 SZ | Anode, Cathode, Sham: Bilaterally over the DLPFC at Fp1 and Fp2 Single reference electrode: right upper arm Stimilation: 1 mA, 20 min, single session EEG and performance-based assessments were obtained at baseline and immediately after stimulation | MMN, P3b (P300) | MMN amplitude: sham > cathodal > anodal group Anodal tDCS significantly decreased MMN amplitudes No effect of tDCS on P300 component was detected |
| Hoy et al., 2015 | Randomized repeated-measures double-blind study | 41 SZ | Anode left DLPFC at F3 Cathode: right supraorbital region Stimilation: 1 mA, 2 mA, and sham, 20 min, single session EEG and performance-based assessments were obtained at baseline and immediately, 20 min, and 40 min after stimulation | 40-Hz gamma synchrony during 2-back working memory task | tDCS with 2 mA significantly improved 40-Hz gamma synchrony at 40 min post-stimulus and 2-back performance at 20 min post-stimulation No change at all measures was detected in other conditions |
| Reinhart et al., 2015 | Randomized cross over study | 19 SZ | Anode medial-frontal cortex at FCz Cathode: right cheek Stimilation: 1.5 mA, 20 min, single session "sham" and "active" tDCS took place at least 48 hours apart | Theta oscillation during color discrimination task | Theta inter-trial coherence (ITC) showed no predictive power over post-error slowing before tDCS application, unlike in HC After active tDCS, peak theta ITC predicted single-trial fluctuation in post-error slowing Anodal tDCS in SZ significantly improved theta ITC and post-error slowing comparable to sham tDCS in HC |
| Reinhart et al., 2015 | Randomized cross over study | 19 SZ | Anode medial-frontal cortex at FCz Cathode: right cheek Stimilation: 1.5 mA, 20 min, single session "sham" and "active" tDCS took place at least 48 hours apart | ERN during color discrimination task | Anodal tDCS in SZ significantly improved ERN amplitude, reaction time, and accuracy comparable to sham tDCS in HC The significant difference in learning dynamics between patients and controls observed at baseline were no longer present after patients received anodal stimulation |
| Author, year   | Study design                  | Patient number | tDCS protocol                                                                 | EEG/ERP measurements                      | Conclusions                                                                                                                                 |
|---------------|-------------------------------|----------------|-------------------------------------------------------------------------------|--------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| Mood disorders |                               |                |                                                                               |                                            |                                                                                                                                            |
| Powell et al., 2014 | Double-blind randomized controlled cross over study | 14 MDD         | Anode: left DLPFC at F3 Cathode: lateral aspect of right orbit at F8 Stimulation: 2 mA, 20 min, single session “sham” and “active” tDCS took place 7–8 days apart | ERS, ERD, P1, N2, P300 during visual working memory task | In active group, significant reduction in the N2 amplitude and theta activity over frontal areas (FCz) during memory retrieval In active group, significant increase in alpha dysnchronization over parietal areas (Pz) during maintenance |
| Al-Kaysi et al., 2016 | Double-blind randomized controlled cross over study | 10 MDD         | Anode: left DLPFC at F3 Cathode: lateral aspect of right orbit at F8 Stimulation: 2 mA, 20 min, 15 sessions for 15 days “sham” and “active” tDCS took place 7–8 days apart | EEG power spectral density | AF8-CZ and AF8-C1 channels were best combinations for classification of MDD patients. Classification results were better for the cognitive improvement (9/10 participants) than for mood improvement (7/10 participants) |
| Al-Kaysi et al., 2017 | Double-blind randomized controlled cross over study | 10 MDD         | Anode: left DLPFC at F3 Cathode: lateral aspect of right orbit at F8 Stimulation: 2 mA, 20 min, 30 sessions over 6 weeks 15 sessions over 3 weeks (double-blind) and additional 15 sessions over additional 3 weeks (open-label) | EEG power spectral density | Improvement in mood following tDCS accurately predicted in 8/10 using FC4-AF8 channel pair (accuracy=76%, p=0.034) Improvement in cognition following tDCS accurately predicted 10/10 using CPz-CP2 channel pair (accuracy=92%, p=0.004) |
| Bersani et al., 2015 | Open label study               | 25 BD          | Anode: left DLPFC at Fp1 Cathode: right cerebellar cortex Stimulation: 2 mA, 20 min, 15 sessions for 15 days | P3a, P3b (P300)                            | No effect of tDCS on P3a component was detected. P3b amplitude baseline < after tDCS treatment P3b latency: baseline > after tDCS treatment |
| Bersani et al., 2017 | Double-blind randomized controlled study | 42 BD          | Anode: left DLPFC at Fp1 Cathode: right cerebellar cortex Stimulation: 2 mA, 20 min, 15 sessions for 15 days | N1, P3a, P3b (P300)                       | No effect of tDCS on N1 amplitude, latency, and P3 amplitude was detected P3b latency decreased significantly on the active group (not in the sham group) |
| Substance use disorders | Randomized cross over study       | 49 AA          | Anode: left DLPFC at F3 Cathode: contralateral supra-deltoid area Stimulation: 1 mA, 10 min, single session “sham” and “active” tDCS took place 7 days apart | P300                                      | In whole alcohol dependence group, P3 mean amplitude decreased for alcohol-related and neutral sounds during tDCS application. After the end of tDCS application, P3 mean amplitude significantly increased for alcohol-related sounds and was not for neutral one Lesch type IV showed increased P3 amplitude at Fz site during and after tDCS application, whereas type II presented decreased P3 amplitude |
| Author, year | Study design | Patient number | tDCS protocol | EEG/ERP measurements | Conclusions |
|--------------|--------------|----------------|---------------|----------------------|-------------|
| da Silva et al., 2013<sup>37</sup> | Randomized controlled study | 13 AA active: 6 sham: 7 | Anode left DLPFC at F3 Cathode: contralateral supra-deltoid area Stimulation 1 mA, 10 min, 5 sessions for 5 days | Cue-reactive N2, P3 CSD during alcohol-related visual go/nogo task | In active tDCS group, depressive symptoms and craving were reduced to a larger extent, and trend for increased change in executive function and for more relapses were present. Active tDCS was able to block the increase in neural activation triggered by alcohol related and neutral cues in PFC. |
| den Uyl et al., 2016<sup>38</sup> | Randomized controlled study | 78 AA (CBM and tDCS) control and sham: 20 active and sham: 19 control and anode: 20 active and anode: 19 | Anode left DLPFC at F3 Cathode: contralateral supra-deltoid area Stimulation: 1 mA, 15 min, 3 sessions for 3 days | Visual P300 during oddball task, during cue-craving task | No change at all ERP measures was detected. |
| Conti et al., 2014b<sup>41</sup> | Randomized controlled study | 13 CA active: 7 sham: 6 | Anode right DLPFC at F4 Cathode: left DLPFC at F3 Stimulation: 2 mA, 20 min, single session | Visual N2 during cue-craving task | ACC activity during N2 segment was similar for neutral images pre- and post-tDCS for both groups. In active group, ACC activity decreased for crack-related cues post-tDCS (opposite effect observed in sham). After single active session, P3 current density in the left DLPFC increased during neutral cues and decreased during crack-related cues (opposite effect observed in sham). After completing 5 active sessions, P3 current density increased in all prefrontal areas during crack-related cues. |
| Conti et al., 2014a<sup>42</sup> | Randomized controlled study | 13 CA active: 7 sham: 6 | Anode right DLPFC at F4 Cathode: left DLPFC at F3 Stimulation: 2 mA, 20 min, 5 sessions for 5 days | Visual P3 during cue-craving task | |
| Nakamura-Palacios et al., 2016<sup>40</sup> | Randomized controlled study | 22 AA, 9 CA AA active/sham: 8/14 CA active/sham: 6/3 | Anode right DLPFC at F4 Cathode: left DLPFC at F3 Stimulation: 2 mA, continuously for 20 min 13 min with 20 min interval in between (13:20:13), 5 sessions for five days | Visual P300 CSD during cue-craving task | In both alcoholics and crack-cocaine users, ventral medial PFC was the brain area with the largest change towards increasing activation under drug-related cues in those subjects that kept abstinence during and after the treatment with bilateral tDCS over DLPFC, applied repetitively. |
| Author, year | Study design | Patient number | tDCS protocol | EEG/ERP measurements | Conclusions |
|-------------|--------------|----------------|---------------|-----------------------|-------------|
| Alzheimer’s disease | | | | | |
| Khedr et al., 2014 | Double-blind randomized controlled study | 34 AD anodal: 11 cathodal: 12 sham: 11 | Anode, Cathode, Sham: left DLPFC at F3 Single reference electrode: contralateral supraorbital region Stimulation: 2 mA, 25 min, 10 sessions for 10 days | P3b (P300) | Both anodal and cathodal tDCS shortened P300 latency. Significant reduction of reaction time shown only after cathodal tDCS |
| Marceglia et al., 2016 | Randomized cross over study | 7 AD | Anode, Cathode: bilaterally over the TPJ Single reference electrode: right deltoid Stimulation: 1.5 mA, 15 min, single session "anodal" and "cathodal" tDCS took place at 7 days apart | Spectral power, inter- and intra-hemispheric coherence | Pre-tDCS, decreased high frequency power was correlated with lower scores of the mini-mental state exam. Post-anodal tDCS, high-frequency power increased in the tempo-parietal area, and increased temporoparietal-occipital coherence correlated with improvement of performance on the word recognition test (WRT). Post-CtDCS, non-specific effect of decreased theta power all over the scalp that was not correlated with the clinical observation at the WRT |
| Developmental disorders | | | | | |
| Pinchuk et al., 2012 | Retrospective analysis | 128 LD 48 mild MR | Anode varied Cathode: ipsilateral mastoid Stimulation: 60–120 mA, 25–45 min, 5–9 sessions with 2–3 days interval | EEG power spectra and frequency spectra | In all groups, EEG parameters disorganization, no or poor frequency and amplitude modulation of rhythms, excess of slow waves, increased amplitude, pathological inter-hemispheric asymmetry, generalized and focal paroxysmal activity were noted. tDCS not only improved neuropsychological test results but also enhanced EEG parameters, showing their approaching the age norms |
| Amatachaya et al., 2015 | Double-blind randomized controlled cross over study | 20 Autism active: 10 sham: 10 | Anode left DLPFC at F3 Cathode: right shoulder Stimulation: 1 mA, 20 min, single session "sham" and "active" tDCS took place 14 days apart | Peak alpha frequency (PAF) | In active group, significant improvements in social and health/behavior domains of autism treatment evaluation checklist (ATEC). PAF significantly increased at the stimulation site, and increased PAF was significantly associated with improvements in the two domains of ATEC impacted by tDCS |
| Cosmo et al., 2015 | Double-blind randomized controlled study | 60 ADHD active: 30 sham: 30 | Anode left DLPFC at F3 Cathode: right DLPFC at F4 Stimulation: 1 mA, 20 min, single session | Functional cortical network (FCN) of the resting EEG | Comparing the weighted node degree within groups pre- and post-tDCS, a statistically significant difference was found in the electrodes located on the target (left frontal) and correlated (occipital, temporal, centroparietal) areas in the active group (no significant results in sham) |

EEG: electroencephalography, ERP: event-related potential, tDCS: transcranial direct current stimulation, MMN: mismatch negativity, SZ: schizophrenia, HC: healthy control, DLPFC: dorsolateral prefrontal cortex, ERN: error-related negativity, ERS: event-related synchronization, ERD: event-related desynchronization, MDD: major depressive disorder, BD: bipolar disorder, AA: alcohol addiction, CSD: current source density, CBM: cognitive bias modification, CA: crack-cocaine addiction, AD: Alzheimer’s disease, TPJ: temporo-parietal junction, LD: learning disorder, MR: mental retardation, ADHD: attention deficit and hyperactivity disorder
lation of EEG/ERPs that reflect such cognitive functions. Although cognitive dysfunction has been considered a major symptom domain of schizophrenia, the direct measurement of cognitive functioning is confounded by factors such as motor speed, psychotic symptoms, and fatigueability. EEG and ERPs can effectively control these confounders and provide more objective evaluation of cognitive functioning with high temporal resolution.25,26 Hoy et al.27 examined whether improvements in working memory following tDCS on the left dorso-lateral prefrontal cortex (DLPFC) would be reflected as changes in 40-Hz gamma synchrony during 2-back working memory task. Authors reported significantly improved working memory performance, as well as a significant decrease in gamma event-related synchronization. Results from this study provided preliminary evidence for enhanced working memory following tDCS, which restored normal gamma oscillatory function in schizophrenia. Reinhart et al.28 demonstrated aberrant oscillatory activity in the frontal cortex to underlie deficits in adaptive control in schizophrenia patients. The effect of anodal tDCS over the medial frontal cortex on improving neurophysiological signature of adaptive control provided causal evidence for executive control. In patients with schizophrenia, tDCS was found to modulate the temporal structure of low-frequency oscillations and synchrony and to improve behavioral response of post-error slowing to a level of healthy comparison group during color discrimination task performance. Using the same tDCS protocol, Reinhart et al.29 also examined whether deficit learning in schizophrenia patients could be improved via increasing error-related negativity (ERN) and demonstrated improved learning rate, as well as the neural signature of prediction error signaling. Results from these randomized cross over studies all together suggested tDCS as a possible treatment option for cognitive enhancement in patients with schizophrenia.

Affective disorders

For studies in affective disorders, 3 studies pertained to major depressive disorder (MDD) and 2 to euthymic bipolar disorder.29-31 In all the 5 studies, anodal tDCS was applied over the left DLPFC, and in the 2 studies in euthymic bipolar patients, cathodal tDCS was additionally applied on the right cerebellar cortex. To examine the effects of applying anodal tDCS over the left DLPFC in MDD patients, Powell et al.29 investigated the N2 ERP component and event-related spectral changes during a visual working memory task performance. In this double blind sham controlled crossover study, intervention dependent effects were found, that in that active tDCS session resulted in significantly reduced N2 amplitude and theta activity over frontal areas during memory retrieval, as well as in significantly increased alpha over occipital-parietal areas during maintenance. These observed effects on EEG components, which are considered to have a source in medial frontal cortex, suggest tDCS to affect cortical functioning beyond focal changes at the stimulation site. Two studies by Al-Kaysi et al. utilized machine learning methods in EEG data of 10 MDD patients to predict clinical outcomes following tDCS applications over the left DLPFC.30,31 The authors demonstrated that a multichannel deep belief network can be used to accurately classify between the EEG data that was recorded after active and sham tDCS and identified features from the baseline resting-state EEG that differentiate patients who respond to tDCS treatment from those who do not.31 Al-Kaysi et al.30,31 also found frontal channels to perform better in predicting clinical scores, while the parietal-occipital channels better in cognition score. These studies demonstrated the feasibility of EEG-based classification that might help better select patients who would further benefit from tDCS treatment in both alleviating depressive symptoms and neurocognitive functioning. Bersani et al. examined the effects of repeated prefrontal-excitatory and cerebellar-inhibitory tDCS on P300 novelty task in euthymic bipolar patients.32,33 P3b reflects working memory update of change while subjects pay attention to stimulus, while P3a is elicited by unexpected stimulus and is thought to reflect automatic reorienting.34 P3 complexes were found associated with higher-order cognitive processing, and their latency suggest the need of cognitive efforts while performing the task.34,35 These two studies by Bersani et al. that physiologically evaluated neurocognitive functioning differed in terms of their sample size (25 vs. 42), designs (open-label vs. double blind placebo-controlled), and electrode positions used (F3 vs. FP1). Nonetheless, both studies consistently reported decreased P3b latency, and their results suggested the potential utility of prefronto-cerebellar tDCS as an inexpensive, convenient, and painless add-on therapy for treating cognitive dysfunctions in euthymic bipolar disorder patients.

Substance use disorders

Three studies examined ERP components in patients with alcohol dependence for effects of anodal tDCS applied over the left DLPFC.36-38 Nakamura-Palacios et al. examined effects of tDCS in 49 alcoholic subjects, who were further subgrouped according to Lesch's typology. Lesch's typology distinguishes 4 subtypes of alcohol-dependent patients based on the patient's family history, personal psychopathology, and theoretical neurobiological background.39 In this randomized cross over study, cognitive and electrophysiological measures demonstrated tDCS-induced frontal activity enhancement specifically in Lesch IV alcoholics, who are associated with pre-morbid cerebral defects, behavioral disorders, and a high social burden.40 For this sub-group, in particular, increased
P3 amplitude during cue-induced auditory paradigm and improved frontal assessment battery scores were reported. Building upon this study, da Silva et al. assessed the effects of repeated tDCS in 13 alcoholic subjects enrolled in a randomized clinical trial. The study reported tDCS effects on craving, ERPs (cue reactivity), and cognitive functions in line with the previous literature, and extended the finding to mood improvement, as well. However, with a trend for more relapses also reported in this study, results all together indicated the need to carefully assess the clinical value of tDCS in future studies. den Uyl et al. investigated effects of tDCS and cognitive bias modification (CBM) in 78 hazardous drinkers. The authors measured not only behavioral indices, such as craving, implicit associations, and approach tendencies, but also visual P300 ERP components elicited by oddball and cue reactivity paradigms. This randomized controlled study demonstrated a specific decrease in cue-induced craving (i.e., not in overall craving), and reported neither behavioral nor electrophysiological effects of repeated CBM and/or tDCS. The remaining three studies examined in in crack-cocaine uses the effects of applying bilateral tDCS with cathodal stimulation over the left DLPFC and anodal stimulation over the right DLPFC. Conti and Nakamura-Palacios investigated tDCS effects on the cue related N2 ERP component of 13 crack-cocaine addicted subjects. This study, applying a single session of prefrontal tDCS was found to specifically modulated anterior cingulate cortex (ACC) activity of N2 during exposure to drug cues in users and indicated tDCS as a promising adjunctive treatment for addiction. The same authors reported the effects for both single and repetitive applications of bilateral tDCS on P3 ERP component during cue reactivity task in 13 crack-cocaine users. This study reported both single and repeated tDCS application to impact cognitive processing of crack-related cues in prefrontal areas as reflected by change in P3 current source density (CSD). Compared to the effect of single dose, however, that of repetitive tDCS extended beyond the DLPFC to the fronto-parietal cortex, orbitofrontal cortex, and ACC. Nakamura-Palacios et al. reported electrophysiologic effects of repetitive bilateral tDCS in 22 alcoholics examined in Klauss et al. study and 9 crack-cocaine users in Conti et al. study. In both alcoholics and crack-cocaine users, the ventral medial prefrontal cortex (VMPFC) was found as the brain area with the largest cue-related P3 CSD change towards increasing activation under drug-related cues in subjects who kept abstinence during and after tDCS treatment. The authors suggested the bilateral DLPFC tDCS to reduce relapses and craving to the drug use and to increase VMPFC activation under drug cues, which may be of a great importance in the control of drug use in drug addiction.

### Alzheimer’s disease

Kheder et al. examined long-term efficacy of repeated cathodal, anodal, or sham tDCS over the left DLPFC in the neurorehabilitation of 34 AD patients. Auditory P300 was used as a biological marker of AD because its latency is known to be pathologically increased in AD, meaning that AD patients have less cognitive reservoir and need more effort in performing working memory update of change. The double-blind randomized clinical trial demonstrated repeated sessions of tDCS, both cathodal and anodal, to not only improve cognitive function, but also reduce the P300 latency. The study’s results suggest that tDCS over the left DLPFC may enhance cognitive functioning in AD patients by modulating P300 mechanism associated with cognitive reservoir and need of effort. In another study, Marceglia et al. investigated effects of bilateral temporo-parietal tDCS on quantitative EEG (QEEG) in 7 AD patients. This study reported anodal tDCS to induce significant increase of high-frequency power in the temporo-parietal area, as well as of temporo-parieto-occipital coherence correlated with improvement in the performance of word recognition test. This finding of partial reversal in the abnormal pattern of EEG activity observed in AD patients following tDCS application suggests tDCS benefits in AD during working memory tasks supported by the modulation of cortical activity.

### Developmental disorders

Three studies examined tDCS induced modulation of resting EEG in children with developmental disorders. Pinchuck et al. conducted retrospective analysis of tDCS effects on neuropsychological measures and QEEG parameters in children with learning disorders and mild intellectual disability. Although tDCS and EEG protocols across subjects varied, due to the study’s retrospective nature, the authors showed tDCS as not only improving neuropsychological test results but also enhancing EEG parameters to approach the age norms. Amatatchaya et al. examined the effects of anodal tDCS over the left DLPFC on peak alpha frequency (PAF), which is known to be significantly decreased in autism, and on clinical symptoms evaluated by autism treatment evaluation checklist. This randomized crossover controlled study demonstrated improvements in social and health/behavior domains of ATEC associated with significant increase in PAF following active tDCS. To explore the potential utility of tDCS in treatment of ADHD, Cosmo et al. assessed the effects of anodal tDCS over the left DLPFC on brain connectivity using the functional cortical network (FCN) based on EEG activity. The study reported increased functional brain connectivity following tDCS, as well as spreading of its modulatory activity.
DISCUSSION

We reviewed 21 articles reporting changes in EEG or ERP measures following tDCS application in psychiatric patients. Our primary aim was to identify electrophysiological markers reflecting the tDCS induced plastic brain changes in association with symptomatic improvement. Such markers can provide valuable information regarding not only the pathophysiology of psychiatric disorders, but also the individualized neuromodulation strategy in psychiatric patients. Subtle change detected by EEG or ERPs may come precede and forecast the later symptomatic improvement, which may not be sufficiently enlarged for detection following a single or small number of tDCS sessions. Furthermore, EEG or ERP has strengths in evaluating the changes in cognitive functioning following tDCS application, because these measures provide more objective biological evaluation of cognitive functioning with high temporal resolution.

In this systematic review, we found that tDCS with EEG or ERPs have been utilized in patient populations of various psychiatric disorders, including schizophrenia, major depressive disorder, euthymic bipolar disorder, alcohol and/or crack-cocaine addiction, AD, learning disorders, mild intellectual disability, autism, attention deficit and hyperactivity disorder. In schizophrenia, ERP components such as MMN, P3b, ERN have been suggested as biomarkers of pathophysiology because they reflect altered neurotransmission, functional connectivity, cognitive dysfunction, as well as clinical symptoms, associated with the disorder.22,23 Thus, these ERP components have been effectively employed in predicting the course of outcome in schizophrenia and even in prodromal psychosis.19,51,52

In line with previous studies, result of this systematic review suggests EEG/ERPs modulated by tDCS to reflect the underlying plastic neural changes that accompany or forecast the forthcoming symptomatic improvement by tDCS in patients with schizophrenia. The findings in patients with MDD that the topological pattern of EEG spectra effectively predict responders of tDCS may aid the selection of clinical population for tDCS application.30,31 In bipolar disorder and AD, impaired P3 complexes reflected specific cognitive dysfunction, such as working memory deficit or pre-attentive processing, evident in the affected patient population. However, the impairments were normalized with cognitive enhancement after tDCS. These findings suggest that plastic neural change by tDCS were reflected in the modulation of EEG/ERPs, that, in turn, consisted improvement of higher-order cognitive functioning. In substance use disorders, altered ERPs were related with substance cue induced craving that normalized with tDCS application, showing that tDCS in addictive disorders may augment substance addiction treatment by modulating neural circuit of craving.37,41-43

Although with varying protocols, tDCS studies in these psychiatric populations employed a region of interest in the prefrontal cortex, especially the left DLPFC. The effect of tDCS on EEG and ERPs were investigated mainly in relations to cognitive functioning in schizophrenia, mood disorders, AD, and developmental disorders, while the focus primarily lied in the cue-induced craving in substance use disorders. Of the 21 studies reviewed, ten studies examined EEG or ERPs changes after a single session of tDCS, 5 studies after no more than 5 sessions, and 6 studies after more than 10 sessions. Irrespective of targeted EEG/ERP parameters or disease entities, 19 studies reported positive results (i.e., changes in EEG/ERPs following tDCS application and/or their association with symptomatic/cognitive improvement), suggesting that even a single tDCS session can effectively modulate brain plasticity, which is reflected by EEG or ERP in psychiatric disorders and may be relevant to the mechanisms underlying symptomatic or cognitive improvement. However, owing to possible publication bias, cautious interpretation is warranted in reading this systematic review.

In strong support of findings in the current systematic review, our recent study in patients with schizophrenia demonstrated tDCS-induced P50 change and associated improvements in auditory hallucination.39 Twice a day, for 5 consecutive weekdays, schizophrenia patients with treatment refractory auditory hallucination received tDCS with anodal electrode on the left DLPFC and cathodal electrode on the left temporo-parietal junction (TPJ). Auditory hallucination symptom severity and P50 sensory gating were assessed before and after the tDCS application. Improvement in auditory hallucination following tDCS was found significantly correlated with tDCS-induced P50 change. Our results indicated P50 sensory gating as a promising marker that may help reveal the mechanisms underlying tDCS effect on auditory hallucination in schizophrenia.

In this systematic review, number of included studies which tested tDCS effect on EEG/ERPs was too small to draw comprehensive conclusions. Each study investigated different EEG/ERPs in different subject population with different tDCS protocols. However, despite such heterogeneity, the included studies strongly support EEG/ERPs to sensitively reflect tDCS-induced brain changes. Taken together, tDCS and electrophysiology may serve useful and sensitive tools for investigating pathophysiology of psychiatric disorders, as well as for selecting regions targeted for neuromodulation in psychiatric disorders. Future studies with consistent tDCS protocols and electrophysiological markers in each disease entities would be strongly suggested to convince and elaborate the findings of studies included in this systematic review.
REFERENCES

1. Wagner T, Valero-Cabre A, Pascual-Leone A. Noninvasive human brain stimulation. Annu Rev Biomed Eng 2007;9:527-565.

2. Brunelini J, Mondino M, Gasab L, Haesebaert F, Gaha L, Suad-Chag- ny MF, et al. Examining transcranial direct-current stimulation (tDCS) as a treatment for hallucinations in schizophrenia. Am J Psychiatry 2012; 169:719-724.

3. Stagg CJ, Nitsche MA. Physiological basis of transcranial direct current stimulation. Neuroscientist 2011;17:37-53.

4. Shiozawa P, Fregni F, Bensenor JM, Lotufo PA, Berlim MT, Daskalakis JZ, et al. Transcranial direct current stimulation for major depression: an updated systematic review and meta-analysis. Int J Neuropsychopharmacol 2014;17:1443-1452.

5. Volpato G, Piccione F, Cavinato M, Duuzzi D, Schiff S, Foscolo L, et al. Modulation of affective symptoms and resting state activity by brain stimulation in a treatment-resistant case of obsessive-compulsive disorder. Neurocase 2013;19:360-370.

6. Aparicio LV, Guariento F, Raza LB, Carvalho AF, Fregni F, Brunoni AR. A systematic review on the acceptability and tolerability of transcranial direct current stimulation treatment in neuropsychiatric trials. Brain Stimul 2016;9:671-681.

7. Brunoni AR, Amadada J, Berbel B, Volz MS, Rizziero BG, Fregni F. A systematic review on reporting and assessment of adverse effects associated with transcranial direct current stimulation. Int J Neuropsychopharmacol 2011;14:1133-1145.

8. Berlim MT, Van den Eynde F, Daskalakis ZJ. Clinical utility of transcranial direct current stimulation (tDCS) for treating major depression: a systematic review and meta-analysis of randomized, double-blind and sham-controlled trials. J Psychiatr Res 2013;47:1-7.

9. Pelletier SJ, Cichetti F. Cellular and molecular mechanisms of action of transcranial direct current stimulation: evidence from in vitro and in vivo models. Int J Neuropsychopharmacol 2014;18.

10. Fritsch B, Reis J, Martinovich K, Schamba HM, Yi Y, Cohen LG, et al. Direct current stimulation promotes BDNF-dependent synaptic plasticity: potential implications for motor learning. Neuron 2010;66:198-204.

11. Cambiaggi M, Velikova S, Gonzalez-Rosa JJ, Cursi M, Comi G, Leocani L. Brain transcranial direct current stimulation modulates motor excitability in mice. Eur J Neurosci 2010;31:704-709.

12. Rissling AJ, Makeig S, Braff DL, Light GA. Neurophysiologic markers of abnormal brain activity in schizophrenia. Curr Psychiatry Rep 2010;12:572-578.

13. Korostenskaja M, Kahlonen S. What do ERPs and ERFs reveal about the effect of antipsychotic treatment on cognition in schizophrenia? Curr Pharm Des 2009;15:2573-2593.

14. Schmidt A, Diaconescu AO, Kometer M, Friston KJ, Stephan KE, Vollenweider FX. Modeling ketamine effects on synaptic plasticity during the mismatch negativity. Cereb Cortex 2013;23:2394-2406.

15. Popov T, Jordanov T, Rockstroh B, Elbert T, Merzenich MM, Miller GA. Specific cognitive training normalizes auditory sensory gating in schizophrenia. Psychiatry Res 2015;228:191-196.

16. Reinhart RM, Zhu J, Park S, Woodman GF. Synchronizing theta oscillations with direct-current stimulation strengthens adaptive control in the human brain. Proc Natl Acad Sci U S A 2015;112:9448-9453.

17. Powell TY, Boonstra TW, Martin DM, Loo CK, Breakspear M. Modulation of cortical activity by transcranial direct current stimulation in patients with affective disorder. PLoS ONE 2014;9:e98503.

18. Al-Kaysi AM, Al-Ani A, Loo CK, Breakspear M, Boonstra TW. Predicting brain stimulation treatment outcomes of depressed patients through the classification of EEG oscillations. Conf Proc IEEE Eng Med Biol Soc 2016;2016:5266-5269.

19. Al-Kaysi AM, Al-Ani A, Loo CK, Powell TY, Martin DM, Breakspear M, et al. Predicting dCPS treatment outcomes of patients with major depressive disorder using automated EEG classification. J Affect Disord 2017;208:597-603.

20. Bersani FS, Minichino A, Fattapposta F, Bernabei L, Spagnoli F, Mannarelli D, et al. Prefrontocerebellar transcranial direct current stimulation increases amplitude and decreases latency of P3b component in patients with euthymic bipolar disorder. Neuropsychiatr Dis Treat 2015;11:2913-2917.

21. Bersani FS, Minichino A, Bernabei L, Spagnoli F, Corrado A, Vergnani L, et al. Predicting positive and negative symptoms response to transcranial direct current stimulation in bipolar patients. J Psychiatr Res 2011;45:785-793.

22. Light GA, Williams LE, Minow F, Sprock J, Rissling A, Sharp R, et al. Electroencephalography (EEG) and event-related potentials (ERPs) with human participants. Curr Protoc Neurosci 2010; Chapter 6: Unit 6. 21-24.

23. Korostenskaja M, Kahlonen S. What do ERPs and ERFs reveal about the effect of antipsychotic treatment on cognition in schizophrenia? Curr Pharm Des 2009;15:2573-2593.

24. Schmidt A, Diaconescu AO, Kometer M, Friston KJ, Stephan KE, Vollenweider FX. Modeling ketamine effects on synaptic plasticity during the mismatch negativity. Cereb Cortex 2013;23:2394-2406.

25. Popov T, Jordanov T, Rockstroh B, Elbert T, Merzenich MM, Miller GA. Specific cognitive training normalizes auditory sensory gating in schizophrenia. Psychiatry Res 2015;228:191-196.

26. Reinhart RM, Zhu J, Park S, Woodman GF. Synchronizing theta oscillations with direct-current stimulation strengthens adaptive control in the human brain. Proc Natl Acad Sci U S A 2015;112:9448-9453.

27. Powell TY, Boonstra TW, Martin DM, Loo CK, Breakspear M. Modulation of cortical activity by transcranial direct current stimulation in patients with affective disorder. PLoS ONE 2014;9:e98503.

28. Al-Kaysi AM, Al-Ani A, Loo CK, Breakspear M, Boonstra TW. Predicting brain stimulation treatment outcomes of depressed patients through the classification of EEG oscillations. Conf Proc IEEE Eng Med Biol Soc 2016;2016:5266-5269.

29. Al-Kaysi AM, Al-Ani A, Loo CK, Powell TY, Martin DM, Breakspear M, et al. Predicting dCPS treatment outcomes of patients with major depressive disorder using automated EEG classification. J Affect Disord 2017;208:597-603.

30. Bersani FS, Minichino A, Fattapposta F, Bernabei L, Spagnoli F, Mannarelli D, et al. Prefrontocerebellar transcranial direct current stimulation increases amplitude and decreases latency of P3b component in patients with euthymic bipolar disorder. Neuropsychiatr Dis Treat 2015;11:2913-2917.

31. Bersani FS, Minichino A, Bernabei L, Spagnoli F, Corrado A, Vergnani L, et al. Predicting positive and negative symptoms response to transcranial direct current stimulation in bipolar patients. J Psychiatr Res 2011;45:785-793.

32. Light GA, Williams LE, Minow F, Sprock J, Rissling A, Sharp R, et al. Electroencephalography (EEG) and event-related potentials (ERPs) with human participants. Curr Protoc Neurosci 2010; Chapter 6: Unit 6. 21-24.

33. Korostenskaja M, Kahlonen S. What do ERPs and ERFs reveal about the effect of antipsychotic treatment on cognition in schizophrenia? Curr Pharm Des 2009;15:2573-2593.

34. Schmidt A, Diaconescu AO, Kometer M, Friston KJ, Stephan KE, Vollenweider FX. Modeling ketamine effects on synaptic plasticity during the mismatch negativity. Cereb Cortex 2013;23:2394-2406.

35. Popov T, Jordanov T, Rockstroh B, Elbert T, Merzenich MM, Miller GA. Specific cognitive training normalizes auditory sensory gating in schizophrenia: a randomized trial. Biol Psychiatry 2011;69:465-471.

36. Reinhart RM, Zhu J, Park S, Woodman GF. Medial-frontal stimulation enhances learning in schizophrenia by restoring prediction error signaling. J Neurosci 2015;35:12232-12240.

37. Knechel L, Schall U, Cooper G, Ramadan S, Stanwell P, Jolly P, et al. Transcranial direct current stimulation of prefrontal cortex: an auditory event-related potential and proton magnetic resonance spectroscopy study. Neurol Psychiatry Brain Res 2014;20:96-101.
Gomes M, de Oliveira RW, de Vasconcellos VF, de Castro LN, et al. Auditory event-related potentials (P3) and cognitive changes induced by frontal direct current stimulation in alcoholics according to Lesch alcoholism typology. Int J Neuropsychopharmacol 2012;15:601-616.

37. da Silva MC, Conti CL, Klauss J, Alves LG, do Nascimento Cavalcante HM, Fregni F, et al. Behavioral effects of transcranial Direct Current Stimulation (tDCS) induced dorsolateral prefrontal cortex plasticity in alcohol dependence. J Physiol Paris 2013;107:493-502.

38. den Uyl TE, Gladwin TE, Wiers RW. Electrophysiological and behavioral effects of combined transcranial direct current stimulation and alcohol approach bias retraining in hazardous drinkers. Alcohol Clin Exp Res 2016;40:2124-2133.

39. Windle M, Scheidt DM. Alcoholic subtypes: are two sufficient? Addiction 2004;99:1508-1519.

40. Lesch OM, Kefer J, Lentner S, Mader R, Marx B, Musalek M, et al. Diagnosis of chronic alcoholism—classificatory problems. Psychopathology 1990;23:88-96.

41. Conti CL, Nakamura-Palacios EM. Bilateral transcranial direct current stimulation over dorsolateral prefrontal cortex changes the drug-cued reactivity in the anterior cingulate cortex of crack-cocaine addicts. Brain Stimul 2014;7:130-132.

42. Conti CL, Moscon JA, Fregni F, Nitsche MA, Nakamura-Palacios EM. Cognitive related electrophysiological changes induced by non-invasive cortical electrical stimulation in crack-cocaine addiction. Int J Neuropsychopharmacol 2014;17:1465-1475.

43. Nakamura-Palacios EM, Lopes IB, Souza RA, Klauss J, Batista EK, Conti CL, et al. Ventral medial prefrontal cortex (vmPFC) as a target of the dorsolateral prefrontal modulation by transcranial direct current stimulation (tDCS) in drug addiction. J Neural Transm (Vienna) 2016;123:1179-1194.

44. Klauss J, Penido Pinheiro LC, Silva Merlo BL, de Almeida Correia Santos G, Fregni F, Nitsche MA, et al. A randomized controlled trial of targeted prefrontal cortex modulation with tDCS in patients with alcohol dependence. Int J Neuropsychopharmacol 2014;17:1793-1803.

45. Khedr EM, Gamal NF, El-Fetoh NA, Khalifa H, Ahmed EM, Ali AM, et al. A double-blind randomized clinical trial on the efficacy of cortical direct current stimulation for the treatment of Alzheimer's disease. Front Aging Neurosci 2014;6:275.

46. Parra MA, Ascencio LL, Urquina HF, Manes F, Ibanez AM, P300 and neuropsychological assessment in mild cognitive impairment and Alzheimer's dementia. Front Neurol 2012;3:172.

47. Marceglia S, Mracic-Sposta S, Rosa M, Ferrucci R, Mameli F, Vergari M, et al. Transcranial direct current stimulation modulates cortical neuronal activity in Alzheimer's disease. Front Neurosci 2016;10:134.

48. Pinchuk D, Vasserman M, Srbiladze K, Pinchuk O. Changes of electrophysiological parameters and neuropsychological characteristics in children with psychiatric development disorders after transcranial direct current stimulation (tDCS). Pol Ann Med 2012;19:9-14.

49. Amatachaya A, Jensen MP, Patjanaasontorn N, Auvichayapat N, Suphakumpinyo C, Janjarasjitt S, et al. The short-term effects of transcranial direct current stimulation on electroencephalography in children with autism: a randomized crossover controlled trial. Behav Neurol 2015;2015:928631.

50. Cosmo C, Ferreira C, Miranda JG, do Rosário RS, Baptista AE,Montoya P, et al. Spreading effect of tDCS in individuals with attention-deficit/hyperactivity disorder as shown by functional cortical networks: a randomized, double-blind, sham-controlled trial. Front Psychiatry 2015;6:111.

51. Kim M, Lee TH, Yoon YB, Lee TY, Kwon JS. Predicting remission in subjects at clinical high risk for psychosis using mismatch negativity. Schizophr Bull 2018;44:575-583.

52. Nieman DH, Ruhrmann S, Dragt S, Soen F, van Tricht MJ, Koelman JH, et al. Psychosis prediction: stratification of risk estimation with information-processing and premorbid functioning variables. Schizophr Bull 2014;40:1482-1490.

53. Kim M, Yoon YB, Lee TH, Lee TY, Kwon JS. The effect of tDCS on auditory hallucination and P50 sensory gating in patients with schizophrenia: a pilot study. Schizophr Res 2018;192:469-470.