Introduction

Plasticity is defined as the ability of the brain to reorganise itself in order to primarily improve the functioning of the brain networks (1). Plasticity enables the modification of neural interaction that outlasts the experimental manipulation. Artificial brain stimulation such as transcranial magnetic stimulation can perturb the cortical and network oscillations (2). The residual effects of TMS are not confined to the stimulated cortex, but may spread across the functionally connected cortical circuits, such as the cortico-cortical and the cortico-thalamic networks (3). This functional connectivity can be explored using electroencephalogram through the modulation of brain oscillatory activity (4).

The hypotheses that suggest a link between the residual effects of TMS and cortical plasticity are due to the ability of TMS to induce changes that outlast the period of stimulation (4). The plasticity-like effect of TMS implies the potential use of this artificial non-invasive magnetic stimulation for basic neurophysiology research as well as for rehabilitation and therapy. It was proposed that TMS interferes with both neuronal and non-neuronal processes (5). The neuronal mechanisms range from local cellular changes to global-scale alteration of neuronal circuits such as network oscillations (6). The cellular changes...
consist of the local synaptic processes of synaptic excitation, synaptic inhibition, and synaptic plasticity, which are akin to the mechanisms of plasticity of long-term potentiation and long-term depression in animal studies (7). Other neuronal changes involve neuromodulators such as dopamine, growth factors such as brain-derived neurotrophic factor (BDNF), and early genes proteins (8). In addition to neuronal effects, TMS also alters non-neuronal processes including cerebral blood flow by changes in blood oxygen level dependent (BOLD) and brain metabolic activity such as glucose metabolism [8]. However, it has always been assumed that the mechanism underlying cortical plasticity is the driving force of sustained TMS aftereffects. Unfortunately, the precise mechanism of TMS-induced cortical plasticity particularly in humans remains elusive. The ability of TMS to emulate the patterns of synaptic plasticity in the hippocampus suggests that TMS can affect synaptic efficacy in the neural network and modulate the cortical and network oscillatory activity (2). However, the cortical and network changes induced by TMS are still relatively unknown. Cortical neurons are largely interconnected and consist of various neural networks ranging from the simple, micro-level interconnections, to the dynamic and complex macro-level networks (5). The neural networks are widely implicated in the process of cortical information coding. Network oscillations through the balance of synchronisation and desynchronisation of neural assemblies are the important mechanisms involved during cortical information transfer (5). However, knowledge on the cortical and network oscillatory activity is still limited.

There is increasing evidence that has demonstrated the link between abnormal electrophysiological properties of network oscillations and the generation of neurological and psychiatric disorders (9). Altered brain rhythms are seen in patients of Parkinson’s disease, schizophrenia, epilepsy, neuropathic pain, tinnitus, migraines, major depression, obsessive-compulsive disorder and psychosis. The term “Thalamocortical dysrhythmia (TCD)” describes abnormal prolonged low-frequency oscillations of delta and theta brain rhythms seen in patients of various neurological and psychiatric disorders (10). Although low-frequency oscillations are normal during slow-wave sleep, prolonged slow oscillations, such as theta rhythms, during awake periods and at rest interrupt the complex dynamic flow of information between the thalamus and cortex, and therefore may produce symptoms of neuropsychiatric illnesses (10). The combined TMS-EEG could have wide applicability in clinical research for characterising disturbances in oscillatory patterns and the altered functional connectivity in neuropsychiatric illnesses [9]. By directly entrain the oscillatory brain rhythms in a control manner, the TMS-EEG can indeed offer exciting possibilities as a diagnostic and therapeutic tool (11). Despite the rise in clinical research exploring the therapeutic potential of TMS, and the evidence of altered brain rhythms in neuropsychiatric patients, knowledge of the precise mechanisms of cortical oscillatory activity after the full range of TMS applications is still lacking.

In this review paper, we discuss the neurophysiological mechanisms of TMS aftereffects on cortical and network oscillations in humans. Focusing on EEG as the direct index of cortical output, the goal was to develop a deeper understanding of the modulation of oscillatory activity through various brain rhythms after non-invasive magnetic stimulation. In particular, we discuss what EEG response patterns may emulate LTP-/LTD-like changes that lead to corresponding changes in brain excitability and also provide information on versatile mechanisms of TMS actions, possibly exploitable in therapy.

**Transcranial Magnetic Stimulation**

In 1985, Professor Anthony Barker and colleagues from the University of Sheffield, UK, introduced transcranial magnetic stimulation (12). TMS is a non-invasive, non-pharmacological neurophysiologic method of delivering electrical stimuli by rapidly changing the magnetic field (12). TMS induces current into the brain without physical contact, as there are no implanted or surface electrodes. Instead, it works by placing an electromagnetic coil that carries pulses of current near the human scalp. The current from the TMS coil will generate an intense but brief magnetic field (up to 2 Tesla that lasts for 100μs) that passes through the scalp, skull, and meninges to the cortical region beneath the coil without attenuation (13). Based on Faraday’s law of electromagnetic induction, the rapidly changing magnetic field will induce an electrical current in the surrounding cortical tissue below the coil. As body tissue is electrically conductive, the ionic current will flow, eliciting nerve depolarisation and action potentials, and will subsequently stimulate the cortical neurons (13).

TMS has several advantages. Firstly, TMS does not generate strong pain because the TMS
The applications of TMS-EEG co-registration can be classified into three categories: inductive—using TMS-EEG as index of brain physiological state in behaviourally silent regions; interactive—using TMS-EEG to investigate the functional and dynamics of the brain; rhythmic—using TMS-EEG to study the generation and functional significance of brain rhythms (21). The inductive approach of TMS-EEG uses TMS-evoked potentials (TEP) recorded over the scalp as markers of the internal state of the brain in behaviourally silent areas. The interactive approach of TMS-EEG investigations involves the application of these combined methods to explore the transient modulation of neuronal networks during task performance (20). This approach is mainly used to identify the cortical area that is involved in a particular task. The rhythmic approach uses TMS-EEG to examine the modulation of oscillatory brain activity by rTMS and the link between specific frequency bands and their functional role (21). The significance of this approach is the potential role of using TMS to transiently modify brain functions by altering brain oscillations, and therefore, it may contribute to the therapeutic strategy of using TMS to reverse abnormal synchronisation in neuropsychiatric disorders (22).
Alteration of Cortical Oscillations Induced By TMS

The first study that demonstrated the ability of magnetic stimulation to alter cortical oscillatory activity was by Paus et al. (23). The authors delivered single-pulse TMS over the sensorimotor cortex at rest and showed increased synchronisation of beta band (15–30Hz) that lasted for several hundred milliseconds. This brief increase of synchronisation reflects the ability of TMS to induce the resetting of oscillations in a “resting” brain (23). A follow-up study by Fuggetta et al. (24) showed that single-pulse TMS applied over M1 at rest induced synchronisation in the alpha and beta bands for 500 ms post magnetic stimulation, and increased linearly with stimulus intensity. As Paus et al. stated previously, the authors concluded that the TMS-induced oscillations were linked to the resetting of the cortical oscillators, instead of the “idling” state of the brain (23,34).

In 2006, Van Der Werf and Paus investigated the oscillatory activity of patients with Parkinson’s disease who underwent partial thalamotomy—unilateral surgery of the ventrolateral nucleus of the thalamus (25). Applying TMS over the intact hemisphere, the authors observed higher synchronisation of beta frequency band in the unoperated hemisphere (with the thalamus intact) than in the operated hemisphere (with thalamotomy) (25). This result implies the role of the thalamus in generating cortical oscillatory activity through various cortico-cortical networks and cortico-thalamic feedback loops (26). However, the oscillating properties depend on the connectivity of different pacemakers and the modulation of the reticular system, which is interconnected with all the thalamic nuclei (26). Besides thalamus, basal ganglia have an important role in driving oscillatory activity in the human motor cortex during motor performance (27). Using TMS, it has been shown that beta frequencies is prominent during tonic contraction but is attenuated prior to and during voluntary movement (28). In Parkinson’s disease, the alterations of basal ganglia physiology may involve the alteration in the pattern of neuronal synchronisation particularly involving beta brain rhythms (28). The level of beta synchronisation is in turn modulated by net dopamine levels at sites of cortical input to basal ganglia (29). Dopamine deficiency as in the case of Parkinson’s disease will disrupt the cortico-basal ganglia-thalamocortical circuits, leading to pathologically exaggerated beta oscillations (29).

In recent years there has been a growing interest in the cortical oscillatory activity at “rest” as an index of the internal state of the brain (30). The term “rest” represents the cortex during behaviourally silent states, with the absence of any sensory or motor output (30). The properties of neuronal oscillatory brain rhythms in a resting brain can provide the baseline for researchers and clinicians in distinguishing the oscillatory patterns that may be disrupted in patients of various neuropsychiatric disorders. However, only few studies have examined the potential use of TMS to transiently modulate brain rhythms over primary motor cortex (M1) at “rest”. These studies have shown that the response of the EEG oscillatory state of the sensorimotor cortex at “rest” depends on TMS intensity, frequency of magnetic stimulation, and the total number of magnetic pulses. Strens et al. (2002) applied a train of 1500 pulses of 1Hz repetitive TMS (rTMS) for 25 minutes over M1 at a subthreshold intensity (31). They demonstrated a decrease in EEG power of α frequency band of 6% and a focal increase of coherence during active task compared to resting condition ipsilateral to the site of stimulation [31]. A follow up study by Oliviero et al. (32) used a short train of 50 pulses of high frequency 5Hz rTMS over M1 at active motor threshold. They showed a significant decrease in cortico-cortical interhemispheric coherence in the upper alpha frequency band (10.7–13.6Hz) between the motor and premotor cortex for a few minutes after magnetic stimulation (32). In an online rTMS-EEG study, Fuggetta et al. (2008) used spectral analysis of event-related power (ERPow) and event-related coherence (ERCoh) to reveal how intermittent short trains of high frequency (5Hz) rTMS delivered over left M1 induced an ERPow increase in upper alpha (10–12Hz) and beta (18–22Hz) frequency ranges for threshold (100% RMT) and subthreshold intensities (80% RMT). ERCoh showed a decrease in functional coupling for subthreshold rTMS in alpha and threshold rTMS for beta band (33). However, the aftereffect of rTMS in this experiment was short lasting—confined to 500ms after the magnetic stimulation—with no effect found two seconds after the train of magnetic pulses (33). Another on-line rTMS-EEG study by Brigioni et al. explored the immediate effects of low frequency 1Hz rTMS on the ongoing cortical oscillatory activity at “rest” (34). They delivered 1Hz rTMS over M1 at 110% AMT of 600 stimuli. They showed a simultaneous increase of synchronisation of α (8–12Hz) more than β (12–30Hz) across all three stimulation blocks, which was inversely
correlated with the progressive decrease of MEP amplitude [34]. Veniero et al. investigated the effects of the ongoing oscillatory activity of M1 at rest after high frequency 20Hz rTMS (35). They observed increased synchronisation in alpha (8–12Hz) more than beta (13–30Hz), and alpha induction lasted for 5-min after magnetic stimulation. They showed a dose dependent increase of synchronisation in both the alpha and beta activities, spreading from the central region to the posterior, parietal sites (35).

In most TMS studies involving humans, low and high stimulation frequencies often result in opposite physiological effects as index by motor evoked potentials (MEP), expressed as either an increase or decrease in the amplitude of MEP (20). Low stimulation frequency (≤ 1Hz) decreases cortical excitability (MEP suppression) whereas high frequency stimulation (≥ 1Hz) increases cortical excitability (MEP enhancement) (17). However, rTMS-EEG studies of low and high frequency protocols were not able to emulate the classical dichotomy between low versus high frequency rTMS of MEP measurements (35). Instead, they observed linear EEG synchronisation for both low and high frequency rTMS in both alpha and beta frequency bands (33, 34, 35). The inability of rTMS-EEG to distinguish the opposite effect of low versus high frequency at the cortical level may be because alpha and beta frequency bands are not the best index to reflect the dichotomy between low versus high frequency. A differential effect of low-high frequency rTMS may be better demonstrated by the modulation of other brain rhythms such as theta oscillations (35).

Noh et al. investigated the short-term modulation of cortical oscillations after high frequency rTMS by manipulating the different number of magnetic pulses (36). They compared the cortical readout of direct electrophysiological EEG after high frequency rTMS (~11Hz) of 20 trains of 20 pulses (400 magnetic pulses; rTMS 20) versus 20 trains of 60 pulses (1600 magnetic pulses; rTMS 60) over the left M1 at rest. They found the distinctly different topography and temporal dynamics of theta and mu rhythms (36). The theta synchronisation was globally distributed across multiple locations of the EEG electrodes for 20 seconds after rTMS 60 pulses. The mu rhythm was focally distributed and dominated early for 5 seconds after rTMS 20 pulses. These findings point to the probable presence of independent theta and mu generators over the human motor network with different reactivity to rTMS (36). Fuggetta and Noh investigated whether low frequency brain rhythms such as delta and theta oscillations could be used to exhibit the dichotomy between the simple protocols of low and high frequency magnetic stimulation (9). Short trains of low frequency 1Hz rTMS versus high frequency 5Hz and 10Hz rTMS over M1 at rest were applied with simultaneous EEG recordings (9). Their results showed the ability of low frequency EEG oscillations of delta and theta brain rhythms to contrast the modulatory effects of low and high frequency rTMS (9). The findings of these experiments on the short-lasting modulation of low frequency oscillations after rTMS suggest that short trains of rTMS were able to induce short-term plasticity-like mechanisms over the motor cortex. Although the rTMS-induced short-term plasticity-like mechanisms are enough for basic neuroscience research, the short-lasting effects are not sufficient for clinical intervention.

In order to investigate whether TMS could modulate EEG oscillatory activity for relatively longer periods of time, a pattern rTMS protocol of continuous theta burst stimulation (cTBS) was used. cTBS has been shown to induce longer-lasting behavioural effects after magnetic stimulation (37). In this experiment, the authors applied 300 pulses of short intensity but high frequency cTBS over the left M1 and measured the EEG oscillatory activity both at rest and during an active motor task [38]. Their results showed that cTBS could modulate the cortical brain rhythms, particularly beta oscillations, for at least 30 minutes compared to the 20 minutes MEP suppression for both event-related power modulation (ERPow) and event-related coherence (ERCoherence) (39). This finding suggests that EEG is probably a more sensitive index of cortical output after cTBS compared to MEP.

**Cortical Plasticity Induced By TMS**

The hypotheses that suggest a link between the residual effects of TMS and plasticity is due to the ability of TMS to induce changes that outlast the period of stimulation (5). The residual effects of TMS are thought to originate from synaptic plasticity because its effects tend to emulate the patterns of synaptic plasticity in the rodent hippocampus (5). The long-term changes in the strength of hippocampal synapses involve the mechanisms of long term potentiation (LTP) and long term depression (LTD), which describe the direction of a long-lasting change in synaptic strength (40). LTP is an increase in the synaptic efficacy that could last for hours, days or weeks following brief high-frequency stimulation.
The cellular basis of LTP and LTD originates from the hippocampal synapses of the axons of CA3 neurons and the dendritic spines of CA1 pyramidal neurons (42). The CA3 axon terminals discharge glutamate while the CA1 neurons express three types of glutamatergic receptors: α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPA-R), N-methyl-D-aspartate receptor (NMDA-R), and metabotropic glutamate receptor (mGluR). The induction of LTP depends on the influx of Ca\(^{2+}\) in the postsynaptic cell. It starts when glutamate binds to AMPA-R, allowing Na\(^{+}\) to enter into the dendritic spine, resulting in membrane depolarisation (42). When the postsynaptic neuron is sufficiently depolarised, the Mg\(^{2+}\) ions that block the NMDA-R at resting membrane potential will be removed, thus opening the NMDA-R (43). As a result, Ca\(^{2+}\) enters the postsynaptic neuron, and activates calcium-sensitive signaling pathways such as calcium-calmodulin protein kinase II (CaMKII) that leads to phosphorylation and upregulation of the AMPA-R. HFS protocols during experimental stimulations are able to remove the Mg\(^{2+}\) block of the NMDA-R, probably because HFS activates many AMPA-R, thus eliciting a large depolarisation in the dendritic spine. The mechanism of LTD induction also depends on NMDA-R activation, which elevates Ca\(^{2+}\) concentration postsynaptically [44]. The element that determines whether LTP or LTD is induced is the nature of the Ca\(^{2+}\) signal that activates specific pathways. Large and fast elevation of Ca\(^{2+}\) concentration induces LTP by activating CaMKII, whereas small and slow rises of Ca\(^{2+}\) induce LTD by activating protein phosphatases that leads to dephosphorylation and down-regulation of the AMPA-R (45). LTD can be induced by LFS protocols that will mildly stimulate NMDA-R and produce an intermediate elevation of Ca\(^{2+}\) concentration (45).

Several studies have attempted to extend the principles of synaptic plasticity in the animal investigations to the TMS alterations of human cortical excitability. Primary motor cortex has been the most investigated cortical region with regards to TMS-induced plasticity. The studies highlight the success of TMS protocols in emulating the induction paradigms for LTP and LTD by changes in MEP sizes that outlast the TMS application. One TMS protocols that is able to produce long-term changes that emulate the protocols used for inducing LTP and LTD in rodent preparations is theta burst stimulation. Huang et al. showed that two TBS modalities have opposite effects on motor cortex excitability reminiscent of LTP and LTD (37). The iTBS of 600 pulses at 80% AMT produce a facilitatory effect for 15 minutes, whereas cTBS of 300 or 600 pulses suppress MEP amplitude for 20 or 60 minutes, respectively (46). Other studies of iTBS and cTBS showed similar results. Several pharmacological studies of TBS in the human cortex revealed that NMDA receptors seem to have parallel roles in the plasticity of cortical synapses as in the hippocampus. In a double-blind placebo-control study, Huang et al. (2007) investigated the residual effects of TBS by prescribing the NMDA-R antagonist memantine and measuring the MEP size (47). The authors discovered that memantine blocked both the facilitatory effect of iTBS and the suppressive effect of cTBS as shown in the difference of the MEP size compared to control. Other recent studies showed that TBS influences NMDA receptor activity in humans and thus provide evidence of the involvement of TMS in neuroplasticity (48).

Although the modulatory changes in cortical efficacy by rTMS seem to emulate the paradigms of synaptic plasticity, it is important to emphasis the difference between the plasticity studies of the animal hippocampus and TMS studies of the cerebral cortex. The excitation of neural tissue in animal studies of synaptic plasticity and TMS studies in humans is fundamentally different (45). The stimulation of hippocampal slices in LTP/LTD studies is focal, whereas rTMS stimulation has a larger spatial resolution ranging from mm\(^2\) to cm\(^2\). Moreover, the brain region stimulated by rTMS in human studies is the cerebral cortex, which has a structurally more complex network than the hippocampal circuits (46). Cortical neurons are placed in multi-layered arrangements (the canonical six layers), with abundant synaptic connections. Cortical neurons receive massive inputs from the thalamus and, in turn, project heavily to the same structure. Therefore, this suggests that TMS may affect the vast recursive loops of excitation and inhibition between the cortex and the thalamus, between the different areas of the cortex, and including loops...
of both cerebral hemispheres (48). Moreover, the majority of rTMS-induced plasticity studies in humans used MEP amplitudes, which represent an indirect index of plasticity at the neuronal level. In addition, MEP is a polysynaptic read-out, separated by at least three synapses from the TMS source, whereas LTP and LTD are monosynaptic events (48). Therefore, in order to obtain a more accurate interpretation, it is important to combine rTMS with a recording technique that is also linked by a single synapse to the TMS pulse. One such technique is high-density EEG, which can provide a monosynaptic cortical readout during and after magnetic stimulation (49).

Studies that combined TBS and EEG to investigate the effect of cortical excitability induced by magnetic stimulation are lacking (50). A study that examined EEG network oscillations post-cTBS of 600 pulses was performed on the frontal eye field of only four healthy subjects (51). The study demonstrated higher neuronal synchronisation of the cerebral hemisphere ipsilateral to the stimulation site relative to the non-stimulated hemisphere up to one hour with synchronisation computed for broadband EEG and all brain rhythms. The authors speculated that cTBS might interfere with information transfer through its effect on neuronal synchronisation (51). However, in their study, there was no direct comparison between surface EEG and behavioural measurements during rest and active conditions to look at post-cTBS cortical plasticity effects. Moreover, the authors changed the site of stimulation (frontal eye field instead of motor cortex), the stimulation intensity (80% RMT instead of 80% AMT), and modified the cTBS paradigm (30 Hz bursts repeated at 6 Hz) from the original cTBS protocol introduced by Huang et al. (2005), making direct comparison with the original protocol problematic (52). McAllister et al. investigated the modulation of cortical oscillatory activity by cTBS of 600 pulses after a visuomotor training task using both MEP and EEG measurements (53). The authors only found significant alpha power that was positively correlated with MEP after the visuomotor training. They concluded that EEG was not useful as an index of cortical output to plasticity-inducing paradigms such as cTBS. However, in that study, the EEG was recorded using a single electrode of C3 over the motor cortex, and was therefore unable to ascertain the possible cTBS effects on cortico-cortical coupling (53). An investigation using multi-channel EEG will provide a more thorough outlook on the effects of cTBS on the motor network excitability.

In a subsequent experiment, Noh et al. addressed the lack of knowledge of cTBS effects on motor network oscillations and their correlation with behavioural measurements by applying the original cTBS protocol consisting of 100 bursts of three pulses (300 pulses) at 50Hz repeated every 200ms (5Hz) in 13 healthy subjects and measured the EEG oscillatory properties using high-density multi-channel EEG (37). Their results showed that cTBS could modulate the cortical brain rhythms, particularly beta oscillations, for at least 30 minutes compared to the 20 minutes MEP suppression. This finding suggests that EEG is probably a more sensitive index of cortical output after cTBS compared to MEP (37).

In order to demonstrate the link between TMS and oscillatory brain dynamics and cortical plasticity, Table 1 summarises the TMS-EEG co-registration studies that explores the neural correlates.
| **rTMS Protocol** | **Study** | **Site** | **Intensity** | **Number of pulses** | **EEG measures** | **Notes** |
|-------------------|-----------|----------|--------------|---------------------|-----------------|----------|
| rTMS 1Hz | Strens et al. (2002) | M1 | 90% AMT | 1500 | ERCoh alpha | Increase corticocortical & interhemispheric coherence in alpha for 25-min |
| TMS 5Hz | Oliviero et al. (2003) | M1 | 90% AMT | 50 | ERCoh alpha | Decrease ipsilateral corticocortical intrahemispheric coherence in upper alpha |
| rTMS 1Hz | Brignani et al. (2008) | M1 | 110% RMT | 600 | ERPow for alpha and beta | Increase ERPow alpha for 10 minutes inversely correlated with MEP |
| rTMS 5Hz | Fuggetta et al. (2008) | M1 | 80% vs. 100% RMT | 400 | ERPow, ERCoh upper alpha and beta | Increase ERPow alpha > beta. Decrease ERCoh, effect < 2 s |
| cTBS | Schindler et al. (2008) | FEF | 80% AMT | 600 (3 pulses at 30Hz, repeated every 100ms) | Spectral power delta, theta, alpha, beta, gamma, for 60-min | EEG synchronisation for stimulated hemisphere relative to non-stimulated hemisphere for all frequency bands |
| rTMS 20Hz | Veniero et al. (2011) | M1 | 100% RMT | 400 | ERPow alpha and beta | Dose dependent increase ERPow alpha > beta for 5 minutes, inversely correlated with MEP |
| rTMS 11Hz | Noh et al. (2011) | M1 | 100% RMT | 400 vs. 1200 | ERPow and ERCoh theta, mu and beta | Increase ERPow theta > mu > beta for 1200 pulses > 400 pulses |
| cTBS | McAllister et al. (2011) | M1 | 80% AMT | 600 (3 pulses at 50Hz, repeated every 200ms) for 40s | Spectral power for baseline delta, theta, alpha, beta after visuo-training task | No increase EEG synchronisation at rest, MEPs suppression, increase EEG power α after visuomotor training task |
| cTBS | Noh et al. (2012) | M1 | 80% AMT | 300 (3 pulses at 50Hz, repeated every 200ms) for 20s | ERD/ERS, ERPow and ERCoh theta, alpha, low beta, high beta at rest and active; MEP at rest; reaction time active | EEG at rest, EEG active for 30-min, MEP at rest, RT active |
| rTMS 1Hz, 5Hz, 10Hz | Noh et al. (2013) | M1 | 100% RMT | 400 | ERPow theta, mu and beta | Increase ERPow theta > mu > beta for 20s at rest |
| rTMS 1Hz, 10Hz | Noh et al. (2015) | M1 | 100% RMT | 400 | ERCoh delta, theta, mu and beta | Increase ERCoh delta > theta > mu > for 20s at rest |
Limitations and Suggestions for Future Research

Although TMS-induced modulatory aftereffects share many similarities with the mechanisms of synaptic plasticity, the evidence for such associations is, however, indirect. Studies of combined TMS or neuroimaging techniques such as PET and fMRI, and electrophysiological techniques such as EEG and MEG, have found strong indirect links between TMS and plasticity, but direct evidence is still lacking. Animal studies can offer better flexibility in order to establish a direct link between rTMS and plasticity.

Moreover, synaptic plasticity is probably not the only mechanism underlying the residual effects of magnetic stimulation because the EEG oscillatory activity is not correlated with MEP amplitude. Although TMS and plasticity share many characteristics such as TMS has effects that outlast the experimental manipulation, the temporal pattern of TMS—the frequency dependency effects—is similar to LTP/LTD. TMS plays a role in learning, TMS directly impairs or facilitates LTP in rats; however, there is no causal proof that the underlying mechanisms of LTP/LTD and TMS are identical. It is more likely that there is a multiplicity of mechanisms driving the sustained TMS aftereffects (54). Alternative mechanisms driving the modulatory aftereffects of rTMS are altered membrane excitability due to the influence of membrane potentials and ion channels (2). The membrane potential is an important determinant of excitability. The response of a nerve to sequences of impulses at sub- or supra-threshold levels results in a time dependent pattern of excitability changes, which follows changing levels of depolarisation and hyperpolarisation at the axonal membrane. Other alternative mechanisms include reduced cortical excitability in the resting states, increased excitability at the spinal cord, and breakdown of cortical inhibition (19). Future studies should address the multiplicity of mechanisms that drive the rTMS aftereffects besides LTP-/LTD-like mechanisms.

The ability of rTMS to modulate low frequency brain rhythms such as theta oscillations is an exciting phenomenon (36). Evidence from EEG and MEG studies demonstrate that the common link among a wide range of neuropsychiatric disorders is the perturbation of the thalamocortical resonance known as Thalamocortical dysrhythmia (TCD) (9,10). The idea behind TCD is that persistent, abnormal, internally generated delta or theta oscillations in the thalamic neurons disrupt the normal, state-dependent, flow of information within the thalamo-cortico-thalamic network. Although the occurrence of low frequency oscillations is normal during slow-wave sleep, during awake periods, and at rest, prolonged delta and theta rhythms may lead to the disturbances of sensation, motor performance and cognition observed in a number of disorders including Parkinson’s disease, schizophrenia, epilepsy, neuropathic pain, tinnitus, major depression, and obsessive-compulsive disorder (55). In parallel, several TMS protocols have been shown to be able to improve symptoms of various neuropsychiatric disorders although the optimal parameters of magnetic stimulation remain elusive (56). However, to our knowledge, there is very limited studies that investigates the probable link between TMS aftereffects and the TCD phenomenon in clinical populations. Can rTMS reverse TCD, thus alleviating the numerous symptoms in neuropsychiatric disorders? Future clinical trials can exploit the ability of combined rTMS-EEG to modulate and measure the dysrhythmic thalamocortical oscillatory activity in neuropsychiatric disorders.

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