Guiding transcranial brain stimulation by EEG/MEG to interact with ongoing brain activity and associated functions

Citation for published version (APA):

Document status and date:
Published: 01/05/2017

DOI:
10.1016/j.clinph.2017.01.003

Document Version:
Publisher's PDF, also known as Version of record

Document license:
Taverne

Please check the document version of this publication:
• A submitted manuscript is the version of the article upon submission and before peer-review. There can be important differences between the submitted version and the official published version of record. People interested in the research are advised to contact the author for the final version of the publication, or visit the DOI to the publisher's website.
• The final author version and the galley proof are versions of the publication after peer review.
• The final published version features the final layout of the paper including the volume, issue and page numbers.

Link to publication

General rights
Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
• You may not further distribute the material or use it for any profit-making activity or commercial gain
• You may freely distribute the URL identifying the publication in the public portal.

If the publication is distributed under the terms of Article 25fa of the Dutch Copyright Act, indicated by the "Taverne" license above, please follow below link for the End User Agreement:
www.umlib.nl/taverne-license

Take down policy
If you believe that this document breaches copyright please contact us at:
repository@maastrichtuniversity.nl
providing details and we will investigate your claim.

Download date: 17 Sep. 2023
Review

Guiding transcranial brain stimulation by EEG/MEG to interact with ongoing brain activity and associated functions: A position paper

Gregor Thut,⇑ Til Ole Bergmann, Flavio Fröhlich, Surjo R. Soekadar, John-Stuart Brittain, Antoni Valero-Cabrè, Alexander T. Sack, Carlo Miniussi, Andrea Antal, Hartwig Roman Siebner, Ulf Ziemann, Christoph S. Herrmann

Centre for Cognitive Neuroimaging, Institute of Neuroscience and Psychology, University of Glasgow, Glasgow, UK
Department of Neurology & Stroke, and Hertie Institute for Clinical Brain Research, Institute for Medical Psychology and Behavioral Neurobiology, University Hospital Tübingen, Eberhard Karls University of Tübingen, Tübingen, Germany
Department of Psychiatry & Department of Biomedical Engineering & Department of Cell Biology and Physiology & Neuroscience Center & Department of Neurology, University of North Carolina at Chapel Hill, Chapel Hill, USA
Applied Neurotechnology Lab, Department of Psychiatry and Psychotherapy & MEG Center, University Hospital of Tübingen, Tübingen, Germany
Nuffield Department of Clinical Neurosciences, Charles Wolfson Neuroscience Clinical Research Facility, John Radcliffe Hospital, University of Oxford, Oxford, UK
Cerebral Dynamics, Plasticity and Rehabilitation Group, Frontlab, Institut du Cerveau et la Moelle (ICM), CNRS UMR 7225-INSERM U-117, Université Pierre et Marie Curie, Paris, France
Department of Cognitive Neuroscience, Faculty of Psychology and Neuroscience, Maastricht University, Maastricht, The Netherlands
Department of Clinical Neurophysiology, University Medical Center, Göttingen, Germany
Danish Research Centre for Magnetic Resonance, Centre for Functional and Diagnostic Imaging and Research, Copenhagen University Hospital Hvidovre, Hvidovre, Denmark
Department of Neurology, Copenhagen University Hospital Bispebjerg, Copenhagen, Denmark
Department of Neurology & Stroke, and Hertie Institute for Clinical Brain Research, University Tübingen, Tübingen, Germany
Experimental Psychology Lab, Department of Psychology, Center for Excellence “Hearing4all”, European Medical School, Carl von Ossietzky University & Research Center Neurosensory Science, University of Oldenburg, Oldenburg, Germany

* Corresponding author at: 58 Hillhead Street, Glasgow G12 8QB, UK.
E-mail address: gregor.thut@glasgow.ac.uk (G. Thut).

We outline the opportunities of timing NTBS to ongoing brain activity for enhancing its efficacy.
Emerging ideas emphasize brain oscillations as promising targets for interventions.
This offers a principled framework for influencing the brain-behavior relationship by NTBS.

Non-invasive transcranial brain stimulation (NTBS) techniques have a wide range of applications but also suffer from a number of limitations mainly related to poor specificity of intervention and variable effect size. These limitations motivated recent efforts to focus on the temporal dimension of NTBS with respect to the ongoing brain activity. Temporal patterns of ongoing neuronal activity, in particular brain oscillations and their fluctuations, can be traced with electro- or magnetoencephalography (EEG/MEG), to guide the timing as well as the stimulation settings of NTBS. These novel, online and offline EEG/MEG-guided NTBS-approaches are tailored to specifically interact with the underlying brain activity. Online EEG/MEG has been used to guide the timing of NTBS (i.e., when to stimulate); by taking into account instantaneous phase or power of oscillatory brain activity, NTBS can be aligned to fluctuations in excitability states. Moreover, offline EEG/MEG recordings prior to interventions can inform researchers and clinicians how to stimulate: by frequency-tuning NTBS to the oscillation of interest, intrinsic brain oscillations can be up- or down-regulated. In this paper, we provide an overview of existing approaches and ideas of EEG/...
1. Introduction

Non-invasive transcranial brain stimulation (NTBS) of the human brain has gained notable popularity over the last three decades. Today, NTBS is widely used for experimental and clinical interventions in both healthy participants and patients. This is partially due to the development of transcranial magnetic stimulation (TMS) (Barker et al., 1985) and the re-discovery of transcranial current stimulation (TCS) protocols, including transcranial direct current stimulation (TDCS) (Nitsche and Paulus, 2000) or transcranial alternating current stimulation (TACS) as a variant (Antal et al., 2008), which are suitable in terms of ethics and safety for use in most individuals, and normally well tolerated by the participants (Rossi et al., 2009; Brunoni et al., 2011; Woods et al., 2016). Applications are multifold (Bergmann et al., 2016), ranging from studies on normal brain organization and reorganization (e.g. Fröhlich et al., 2015; Kuo and Nitsche, 2015; Prehn and Flöel, 2015) to biomarking (e.g. by recording TMS-evoked responses in electromyography or EEG, Bortolotto et al., 2015) and the development of plasticity inducing protocols (e.g. Liew et al., 2014; Karabanov et al., 2015; Wessel et al., 2015).

While NTBS has greatly advanced clinical neurophysiology and human neuroscience, some important limitations have become increasingly apparent over the years. One of the main limitations is a lack in understanding how NTBS interacts with brain activity at the neuronal level to give rise to behavioral effects, how these effects can most efficiently be optimized and how they may be influenced by intra- and inter-individual factors. Recent findings indicate a high inter- and intra-individual variability in NTBS outcomes across studies, despite the use of identical protocols, which depends on some known variables (such as age, gender, skull shape and structure, emotional and physiological state of participants before and during stimulation, see e.g. Li et al., 2015; Opitz et al., 2015; Ziemann and Siebner, 2015). However, these variables can only explain a small portion of the altogether considerable variability. Essentially, after-effects of commonly used NTBS protocols are challenging to interpret because of high intra- and inter-individual variability, small effect sizes at the group level, and limited reproducibility (Lally et al., 2013; Horvath et al., 2015). Currently, neurophysiologically grounded models of how NTBS interacts most efficiently with functionally relevant brain activity are largely lacking. A hypothesis-driven approach based on physiologically underpinned models is needed to guide the selection of NTBS parameters among the many to choose from.

The challenge is to understand how to target NTBS in order to efficiently interact with neuronal processes that underlie brain function such as perception, attention, memory, cognition or motor control. One important dimension of targeting is functional and structural neuroanatomy and an important tool is neuronavigation. However, the regional spatial specificity of NTBS is of concern. While the spatial resolution of TMS is relatively good (O’Shea and Walsh, 2007, Bolognini and Ro, 2010, but see Schmidt et al., 2015 for physical variability), TMS induced brain activity may spread from the area under the coil along neuronal connections to associated regions depending on the intensity of stimulation (see e.g. Siebner et al., 2000, 2001; Sack et al., 2007; Bestmann and Feredoes, 2013; Martin-Trias et al., 2016). Although the axonal and transynaptic spread of excitation will be restricted by the anatomically predefined connectivity pattern, the spread of excitation limits inferences as to the anatomical origins and substrates of the associated behavioral effects. The spatial specificity of TCS is even more limited than TMS, due to low spatial resolution (Fertonani and Miniussi, 2016), although modeling of current distribution and new ideas for electrode montages (Miniussi et al., 2013; Kloster et al., 2016) suggest that spatial specificity may be increased to stimulate selective cortical structures in the future (Wang et al., 2014). Importantly, functional brain activity is not only defined by spatially distinct networks, but also by dynamic interactions within and across large-scale network components, reflected in brain oscillations across different frequency bands (Buzsáki and Draguhn, 2004). As a consequence, a recent line of research focused on the promise of adding a temporal to the spatial dimension of targeting, and considering brain oscillations as targets for intervention (e.g. Romei et al., 2016).
In this paper, we outline the opportunities afforded by considering the timing of NTBS intervention relative to ongoing brain activity as a window for enhancing its efficacy. In particular, we point out new ideas for tuning the timing of NTBS to ongoing brain oscillations that are based on recent advances in the understanding of the EEG/MEG signal, and illustrate these ideas with recent evidence from TMS-EEG and/or TCS-EEG/MEG studies. We outline the rationale and promise of this approach (Section 2), which covers new ideas about how to tailor TMS/TCS to promising brain signatures (or motifs) for interventions. We survey the growing body of evidence that this approach might work (Section 3), and consider practical issues on how to best document the effects (Section 4). We consider possible underlying mechanisms suggested by models and empirical data (Section 5), and outline important future lines of research on how to optimize the approach (Section 6). The latter includes the use of real-time interventions and the generation of biologically plausible models to guide the choices of stimulation parameters. Finally, we consider the promise of multimodal neuroimaging (TMS-EEG-fMRI) in future research (Section 7). Fig. 1 illustrates the main ideas of the approach.

2. Rationale of temporally guiding NTBS by oscillatory brain activity

Recent research combining EEG or MEG with TMS or TCS has revealed that oscillatory network activity is a promising neural target for NTBS-based interventions to shape brain-behavior relationships.

In terms of their generation, most brain oscillations are reflecting network activity, as they are generated by specific local or large-scale neuronal networks (Buzsaki and Draguhn, 2004), although local pacemaker cells with an intrinsic rhythm do exist (e.g., in the thalamus), and depend on the current vigilance state as they are under strong control of brainstem neuromodulatory systems (Lee and Dan, 2012; Zagha and McCormick, 2014). In addition, most neuronal oscillations, such as those in the theta (4–8 Hz), alpha (8–12 Hz), beta (15–30 Hz) and gamma bands (>30 Hz) rely to some degree on the phasic inhibitory activity of GABAergic interneurons (Wang, 2010), either expressed at a local level (as for gamma) or at a larger scale (such as for the thalamo-cortical alpha rhythm, see e.g. Lorincz et al., 2009). In contrast, the neocortical slow oscillation (<1 Hz) observed during non-rapid eye movement (NREM) sleep may merely result from spontaneous mini-EPSPs triggering persistent sodium currents (rather than from a pronounced GABAergic drive), which initiate a transient depolarization phase (the ‘up-state’) that turns into a subsequent hyperpolarization phase (the ‘down-state’) due to depolarization-activated potassium currents and synaptic depression (Bazhenov et al., 2002; Hill and Tononi, 2005). By means of its alternating ‘down-states’ of wide-spread neuronal silence and ‘up-states’ of increased neuronal firing, the slow oscillation is grouping faster activity such as sleep spindles (12–15 Hz) (Steriade et al., 1993; Steriade, 2006). Even slower, so-called ‘infra-slow’ oscillations (~0.1 Hz), modulate the amplitude of basically all faster oscillations during wakefulness through yet unknown (maybe phasic neuromodulatory) mechanisms (Monto et al., 2008).

Importantly, brain oscillations are associated with various different ‘circuit motives’ that are recurrent throughout the brain, serving comparable computational functions (Womelsdorf and Everling, 2015). Irrespective of the mechanism of generation, the interplay of excitatory and inhibitory neurons within those circuits usually results in periodic fluctuations of the excitation-inhibition balance (EIB) and eventually in variations in the neurons'
membrane potential between states of relative de- and hyperpolarization (Buzsáki and Draguhn, 2004; Schroeder and Lakatos, 2009). The associated alternation between high and low excitability states at the level of a given neuron results in rhythmic gain modulation of both its synaptic input (with EPSPs more easily overcoming firing threshold) and its output (firing rate being modulated by the neuron’s membrane potential) (e.g., Haegens et al., 2011). This rhythmic input/output gain modulation in turn underlies a series of higher order functional principles of neuronal oscillations, such as communication-through-coherence (Fries, 2005, 2015), hierarchical nesting as indexed by phase-power-coupling (Jensen and Colgin, 2007; Schroeder et al., 2008), phase precession and phase coding (Lisman, 2005; Schyns et al., 2011; Jensen et al., 2014), and gating by pulsed inhibition (Jensen and Mazaheri, 2010). Moreover, disinhibition, reflected in a transient deflection of the EIB towards relative excitation by release from inhibition, has been proposed as an important mechanism serving plastic processes at the network level (Letzkus et al., 2015; Cash et al., 2016).

Together, these principles presumably provide a temporal framework as well as the basic computational building blocks of neuronal network interactions in a variety of sensorimotor and cognitive processes (Varela et al., 2001; VanKullen and Koch, 2003; Buzsáki and Draguhn, 2004; Schroeder and Lakatos, 2009).

When synchronized across sufficiently large populations of interconnected neurons, brain oscillations are observable in the local field potential (LFP) and ultimately also non-invasively in surface EEG and MEG, thereby reflecting instantaneous markers of neuronal network excitability (Buzsáki et al., 2012). Given that a neuron’s current state of excitability is an essential factor modulating both the gating and communication of signals as well as the induction of synaptic plasticity within neuronal networks and given the proposed roles of brain oscillations in a variety of cognitive functions, both the oscillatory phase (reflecting current excitability) and amplitude (reflecting current degree of local neuronal synchronization) represent worthwhile targets for NTBS interventions (see also Thut et al., 2012). For instance, tuning NTBS to high excitability phases of oscillatory brain activity may enhance efficacy of NTBS as compared to when applied at random phases. Similarly, synchronizing or desynchronizing brain oscillations by frequency-tuning of electromagnetic stimulation to ongoing oscillations (e.g. by their entrainment) may offer the opportunity to intervene with brain activity and associated functions at a fundamental (mechanistic) level of network interactions.

3. What is the empirical support that tuning NTBS to oscillatory brain activity works?

Several ideas of how NTBS can interact with neuronal oscillations have emerged. The main distinction is between research on immediate and longer-lasting changes, respectively focusing on the effects during NTBS (resulting from direct neuronal excitation/inhibition and interaction with ongoing brain activity) or the after-effects (due to NTBS-induced longer-term changes in excitability or activity immediately following NTBS and beyond). These can be further subdivided in approaches that (1) trigger TMS/TCS by instantaneous oscillatory phase and/or power, (2) tune TMS/TCS to the natural frequency of the underlying oscillation versus (3) a combination of both, as is outlined below (and is schematically represented in Fig. 1C).

3.1. Enhancing NTBS efficacy by triggering TMS/TCS by instantaneous phase and/or power of underlying brain oscillations

The general idea is that the effectiveness of NTBS can be enhanced by timing NTBS to specific phase and/or power values of ongoing brain oscillations (Fig. 1C, left panel). There are indeed several examples of early EEG-TMS studies which demonstrated – using post hoc trial sorting - a relationship between the effectiveness of a single TMS pulse and the power and phase of ongoing brain oscillations at the time of its delivery, as revealed by phase- and/or power-modulation of the amplitude of motor evoked potentials (MEPs) or phosphene induction to TMS over the motor or visual cortex, respectively. The size of the TMS-evoked MEP scales with the power of ongoing sensorimotor µ-rhythms (8–15 Hz) directly preceding the TMS pulse (e.g., Sauseng et al., 2009a; Schulz et al., 2014). Likewise, phosphene reports depend on the power of posterior alpha oscillations immediately preceding occipital TMS (Romei et al., 2008a,b). Moreover, MEPs and phosphene reports have been shown to vary, respectively, with the instantaneous phase of sensorimotor µ -rhythms (Triesch et al., 2015) and posterior alpha oscillations (Dugue et al., 2011) at time of stimulation. Finally, using EEG-triggered TMS, Bergmann et al. (2012a) explicitly targeted up- and downstates of slow oscillation during NREM sleep. It was shown that motor cortical excitability during deep sleep fluctuates in a phase-dependent manner, with larger MEPs and TMS evoked potentials (TEPs) being evoked during slow oscillation EEG upstates and smaller MEPs/TEPs during slow oscillation downstates of the stimulated motor cortex, and the absolute voltage at the time of stimulation further predicting within-state MEP/TEP amplitude (illustrated in Fig. 2).

In addition, recent studies indicate that power and phase at time of TMS may not only influence the immediate effects of TMS, but also TMS after-effects. It has been suggested that by tuning TMS pulses of plasticity-inducing protocols to instantaneous periods of low versus high excitability, plasticity effects may be enhanced. For example, repeated stimulation into either the low or high excitability phase of the sensorimotor µ-oscillation may induce LTP- and LTD-like motor cortical plasticity respectively (Triesch et al., 2015; Zrenner et al., 2015, 2016), in analogy to theta phase-specific plasticity demonstrated in the rodent hippocampus (Huerta and Lisman, 1993, 1995). The rationale behind this approach is to repetitively generate neuronal input, precisely timed to phases of high excitability/disinhibition, thus increasing the chances of TMS-induced postsynaptic firing, and by extension of spike timing dependent plasticity (STDP)-like processes to occur (Artola et al., 1990; Sjöström et al., 2001). In other words, the oscillatory amplitude and/or phase are used as target windows for plasticity inducing protocols. This requires the repeated, temporally precise targeting to a priori defined periods of high excitability in specific oscillatory frequency bands, which has now become feasible with real-time EEG-triggered TMS (see Section 6.1).

3.2. Targeting brain activity and associated functions by frequency tuning of NTBS to underlying brain oscillations

While the approach depicted above utilizes EEG data on instantaneous phase or power for triggering of TMS to enhance TMS efficacy, an alternative methodology aims at tuning rhythmic stimulation protocols (such as repetitive TMS/rTMS or transcranial alternating current stimulation/TACS) to the frequency of an ongoing brain oscillation (Fig. 1C, middle panel). It has been shown that this protocol can result in phase-coupling (also termed entrainment) between oscillatory brain activity and the external electromagnetic stimulus, opening new ways to investigate (and modulate) the relationship between aspects of oscillatory brain activity (such as its phase and amplitude) and behavior.

In terms of immediate effects during NTBS, there are multiple examples that this approach may indeed lend itself for a targeted intervention into oscillatory brain activity through entrainment that can affect brain function and behavior, both using TMS or
TACS (for TMS examples see, Klimesch et al., 2003; Sauseng et al., 2009b; Romei et al., 2010, 2015; Thut et al., 2011; Chanes et al., 2013, 2015; Hanslmayr et al., 2014; Jaegle and Ro, 2014; Quentin et al., 2015a,b; for TACS examples see, Pogosyan et al., 2009; Feurra et al., 2011; Joundi et al., 2012; Neuling et al., 2012; Santarosca et al., 2014; Helfrich et al., 2014b; Cecere et al., 2015; Witkowski et al., 2016; Chander et al., 2016; Ruhnau et al., 2016; Guerra et al., 2016). However, most of the evidence for the existence of entrainment effects comes from behavioral studies. In contrast, only very few TMS or TACS studies so far have managed to simultaneously record EEG/MEG (due to contamination of the recorded neurophysiological signals by stimulation-induced artifacts). Even fewer have combined the two, i.e. recorded EEG/MEG and documented the associated behavioral effects. However, online registration of the EEG/MEG signal is required to verify NTBS interaction with brain oscillations (here entrainment) as the basis of the behavioral change (see Section 4).

Some of the behavioral studies indirectly supporting entrainment by frequency-tuned TACS show performance measures in specific tasks (e.g. sensory detection) to co-cycle with the applied rhythmic electromagnetic force (e.g. over sensory areas) (Neuling et al., 2012). Similarly, TMS-probed excitability in intracortical circuits as inferred from paired-pulse designs (Hallett, 2007) shows modulation by TACS in a frequency- and phase-specific manner, both in terms of intracortical facilitation (ICF) and short-interval intracortical inhibition (SICI) (Guerra et al., 2016). Some level of entrainment seems to have modulated task performance and cortical excitability in-line with TACS phase, as suggested by these behavioral data. Other behavioral studies revealed that when rTMS or TACS is frequency-tuned to known, task-related oscillations, associated behavioral performance measures are biased in expected directions, i.e. in line with known correlative brain-behavior relationships (Klimesch et al., 2003; Sauseng et al., 2009b; Romei et al., 2010; Hanslmayr et al., 2014; Chanes et al., 2013, 2015; Quentin et al., 2015a; Pogosyan et al., 2009; Joundi et al., 2016).
et al., 2012), suggesting that rTMS or TACS has interacted selectively with the target oscillations and associated function by synchronization.

Using concurrent EEG, others have managed to demonstrate entrainment of brain oscillations during frequency-tuned rTMS. For instance, entrainment of parietal alpha oscillations has been demonstrated during short bursts of alpha-rTMS (5 pulses at individual alpha frequency) targeting the right intraparietal sulcus (IPS) (Thut et al., 2011), and entrainment of prefrontal beta oscillations was observed for a few cycles briefly after the end of stimulation when targeting the left inferior frontal gyrus (IFG) (Hanslmayr et al., 2014). Entrainment during short-burst rTMS is frequency-specific, as reflected in stronger entrainment for stimulation at individual frequencies than flanker frequencies (so far tested for beta-rTMS over motor cortex, see Romei et al., 2015). But are TMS-evoked oscillations actually generated by the same neuronal circuits as the targeted spontaneous oscillations? Support for this assumption comes from recent work demonstrating that alpha oscillations evoked by single-pulse TMS of the visual cortex were modulated by top-down attention in the same direction as spontaneous alpha oscillations, namely increasing in amplitude when visual attention was low and decreasing when it was high, which is opposite to the direction e.g. visual evoked potentials would be modulated (Herring et al., 2015). These studies, therefore more firmly establish entrainment of natural brain oscillations as a possible mechanism underlying the above described behavioral effects (see also Section 5).

For the case of TACS, there is clear electrophysiological evidence from animal work that entrainment is possible (Fröhlich and McCormick, 2010; Ozen et al., 2010), but also evidence for transcranial entrainment in humans is accumulating: Concurrent TACS-EEG data suggests that TACS is able to entrain occipital alpha oscillations, although sophisticated TACS-artifact removal procedures are required to extract the brain signals (Helfrich et al.,

---

**Fig. 3.** Tuning NTBS to frequency of underlying brain oscillations. (A) Entrainment of brain oscillations by rTMS (A1) and TACS (A2) when stimulation is directed to posterior alpha oscillations. (B) Functional consequences in terms of perception of these interventions (B1 and B2). A1. Short bursts of alpha-rTMS over right parietal cortex promotes right parietal alpha-oscillations (relative to sham rTMS), and B1. biases visual perception away from the contralateral to the ipsilateral visual field (relative to rTMS at control “flanker” frequencies). A2. Alpha-TACS entrains occipital alpha oscillations (relative to pre and post EEG measures), and B2. causes visual perception to co-cycle with the entrained alpha rhythm. Reproduced from Thut et al. (2011), Romei et al. (2010) and Helfrich et al. (2014b) with permission.
2014b); but see Noury et al. (2016)). Pioneering works on concurrent TCS-MEG demonstrated the feasibility of recording MEG online to TCS, which (in contrast to TCS-EEG) allows to record oscillations directly from the target brain region underneath the stimulation electrode (Soekadar et al., 2013a; Neuling et al., 2015; Ruhnau et al., 2016). After it was shown that monosinusoidal TACS-EEG/MEG recording is accompanied by various stimulation- and heartbeat-related artifacts (Noury et al., 2016) that are difficult to remove by any established methods (Marshall et al., 2016; Noury et al., 2016), a stimulation protocol was recently introduced that avoids the previously described artifact problems by using an amplitude-modulated TACS signal (Witkowski et al., 2016). It was shown that this protocol could entrain prefrontal midline theta oscillations affecting working memory performance and task-dependent theta power-regulation (Chander et al., 2016).

Together with the above described frequency-specific effects on behavior, these EEG/MEG data are suggestive of the possibility to control oscillatory activity and associated performance measures by frequency-tuned interventions. Examples of studies on frequency-tuning rTMS/TACS to brain oscillations are provided in Fig. 3, including evidence for entrainment in EEG (Fig. 3A) and for meaningful behavioral changes resulting from these interventions (Fig. 3B).

In addition to these immediate effects of NTBS, after-effects have been reported with frequency-specific TACS as well as frequency-specific rTMS. For example, entrainment of spontaneous alpha oscillations via TACS (Herrmann et al., 2012) may result in subsequent increases in alpha power (Zaehle et al., 2010; Neuling et al., 2013; Vossen et al., 2015), a phenomenon, which may indirectly rely on STDP induction in the specific alpha-generating circuits by entrainment, but is not a direct sign of entrainment itself (Zaehle et al., 2010; Vossen et al., 2015, see also Veniero et al. (2015)).

3.3. Combinations of frequency-tuned (3.2) and phase-triggered (3.1) interventions

It is likely that the above frequency- and phase-tuned interventions may be potentiated when combined (Fig. 1C, right panel). One potentially effective variant of this combination is to lock frequency-tuning to specific oscillatory phase angles or power values of ongoing oscillations (using real-time NTBS-EEG/MEG approaches). Indeed, the strength of entrainment of parietal alpha oscillations during frequency-tuned rTMS depends on the alpha-phase at which the rTMS-train catches the ongoing alpha-oscillation, as revealed by post hoc trial sorting (Thut et al., 2011). Similarly, strength of alpha-entrainment during alpha-TACS seems to depend on the ongoing alpha-power (eyes-open vs. closed) (Ruhnau et al., 2016). Although real-time power- or phase-dependent frequency-tuning will likely be advantageous, no study has implemented this approach so far.

A second variant of combining frequency- and phase-tuning is to entrain brain oscillations with frequency-tuned interventions (e.g., TACS), together with the presentation of discrete events (e.g. single TMS pulses, gamma bursts, etc.) at specific NTBS phase angles. The feasibility of this approach has recently been shown for TACS-TMS over motor cortex (Raco et al., 2016). Others have combined two TACS waveforms (Alekseichuk et al., 2016) to emulate the circuit motif of cross-frequency phase-power couplings, reported in many EEG/MEG-studies (see Section 2 above). To this end, Alekseichuk et al. (2016) applied TACS over frontal areas in a cross-frequency regime, while participants were performing a working memory task. Fast gamma-TACS stimulation signals were superimposed on a slower, background theta-TACS oscillation (Fig. 4A), which led to marked changes in working memory performance (Fig. 4B) and brain connectivity (Fig. 4C), depending on the phase gamma-TACS was locked to. Importantly, this was not observed with the gamma-TACS bursts just repeated at theta rate (see Fig. 4).

Other combinations are conceivable, e.g., testing whether the pulses of plasticity inducing TMS protocols would be more effective if tuned to specific phases of simultaneously applied TACS, in particular when the latter is frequency-tuned to physiologically meaningful oscillations (see Goldsworthy et al., 2016). This is analogous to the idea that the efficacy of these protocols may be enhanced if TMS pulses are phase-locked in real-time to EEG-signals (see Section 3.1 above).

A third, related approach relies on the simultaneous, frequency-tuned intervention with two nodes of a network (e.g. by double-site TACS), combined with phase-alignment of these two interventions. The approach is about phase-coupling or phase-decoupling of the two, spatially separated TACS stimulation signals to potentially promote or suppress communication between the two stimulated nodes of the network (in alignment with the principle of communication-through-coherence (Fries, 2005), see also Section 2). Examples can be found in Polania et al. (2012) and Helfrich et al. (2014a). These studies tuned TACS frequencies to the natural rhythm of the network under study, e.g. to theta of a fronto-parietal network (Polania et al., 2012) or gamma of a bi-hemispheric occipital network (Helfrich et al., 2014a). The results suggest that the phase-lag between the two TACS waveforms (in-phase versus out-of-phase) affects the associated functions (i.e., working memory or perception of horizontal motion)1 in line with the notion of interfering with functional connectivity by network coupling/decoupling via double-site TACS.

4. Methodological considerations: proper documentation of effects

The EEG/MEG-informed NTBS approach outlined above relies on principled ideas about the relevance of intrinsic brain oscillations in shaping brain function, and how to interact with them. More specifically, both spontaneously fluctuating and NTBS-regulated brain oscillations are thought to represent modulators of the behavioral outcomes of NTBS, when appropriate timing and frequency of NTBS relative to the oscillations are used. Accordingly, it is important to document not only the behavioral outcome but also the hypothesized electrophysiological underpinnings of this approach, which requires the recording of EEG or MEG simultaneously to the intervention. For instance, for phase-tuned intervention, it is important to verify proper phase-targeting in EEG/MEG, while for frequency-tuned intervention, entrainment should ideally be demonstrated, alongside the behavioral effects. However, there are important challenges in the documentation of the electrophysiological underpinnings of these effects, depending on the chosen protocol (e.g. TMS vs. TACS) and effects of interest (immediate vs. after-effects), mainly due to NTBS-induced artifacts in the EEG/MEG recordings as well as EEG/MEG contaminations due to NTBS-associated peripheral sensations. Below we outline these

---

1 It should be noted that studies using a single “return” electrode (e.g., vertex) for two “active” electrodes (e.g., frontal and parietal) to produce in-/out-of-phase conditions (e.g., Polania et al., 2012) not only vary the phase-lag between the two active sites, but also vary the direction of current flow in the brain tissue: The current flow is fronto-parietal (and vice versa) when both sites are out-of-phase (since in that case the two sites have opposite polarity), but fronto-vertex and parieto-vertex (and vice versa) when the two active sites are in-phase (i.e., same polarity) but out-of-phase with the vertex (i.e. different polarity). This problem is circumvented to a certain degree with local center-surround montages (e.g., Helfrich et al., 2014a), where it can at least be assumed that less current flows between the two local montages. For other, useful electrode options in this regard, see also Bortolotto et al. (2016).
challenges and the experimental designs that allow controlling for them, which are critical for evaluating the success of the approach.

4.1. Documentation of behavioral effects

The assessment of behavioral effects is important in the first place, as these effects are the primary outcome measures of most experimental and clinical interventions. The experimental designs should be chosen to allow testing the benefit of adding a temporal to the spatial dimension of targeting. This can be achieved by implementing appropriate control conditions mimicking conventional approaches, e.g., phase jittering stimulation or use of arbitrary (but non-harmonic) stimulation frequencies relative to the target oscillation.

4.2. Documentation of electrophysiological underpinnings

Documentation of online interactions of NTBS with brain oscillations as the origin of the behavioral effects is problematical because of the NTBS induced electrical artifacts in EEG/MEG online to stimulation. For TMS, these consist of brief but high amplitude deflection in the EEG which can be minimized by using appropriate hardware (Virtanen et al., 1999; Veniero et al., 2009) and further reduced by additional, post hoc artifact reduction procedures (Siebner et al., 2009; Ilmoniemi and Kicic, 2010; Vernet and Thut, 2014). For TACS, the electrical artifact is likewise of high amplitude (reaching mV levels relative to μV neuronal signals) but, in addition, is present continuously, which renders it more resistant to elimination (Noury et al., 2016). On top of these electrical artifacts, each technique is associated with a set of unwanted peripheral sensations. For TMS, these consist of auditory and tactile sensations (and associated cranial muscle potentials) (Nikouline et al., 1999; Mutanen et al., 2013; Rogasch et al., 2014). For TACS, the main physiological contaminations are visual sensations originating from stimulation of the retina (Schwiedrzik, 2009; Schutter and Hortensius, 2010; Schutter, 2016; Laakso and Hirata, 2013), which are frequency-dependent (Turi et al., 2013) and occur with many electrode montages due to the retina’s high sensitivity to

---

Fig. 4. Combined frequency-tuning and phase-triggering. (A) Design: Prefrontal cortex was stimulated in nine TACS conditions, including gamma-TACS bursts nested in theta-TACS cycles (i.e., a cross-frequency phase-power TACS protocol), while EEG and working memory performance was recorded. (B) Theta-gamma TACS enhanced working memory performance. This effect depended on the timing of the gamma-bursts relative to the theta cycle (phase modulation, upper bar plot), as well as on the frequency of the gamma bursts (frequency modulation, middle bar plot) and could not be explained by gamma-burst stimulation simply repeated at a theta-rate without the presence of a theta TACS waveform (DC offset controls, lower bar plot). (C) Prefrontal theta-gamma TACS enhanced global brain connectivity, relative to all other conditions (here illustrated for sham). Reproduced from Alekseichuk et al. (2016) with permission.
Electrical artifacts reduction algorithms have been proposed for TMS-EEG (Ilmoniemi and Kicic, 2010; Vernet and Thut, 2014, see also Rogasch et al., 2016: https://nigelrogasch.github.io/TESA and Herring et al., 2015: www.fieldtriptoolbox.org/tutorial/tms-eeeg for removal pipelines implemented in EEGlab and FieldTrip). Likewise, such algorithms have been implemented for TACS-EEG (Helfrich et al., 2014b, for limitations see Helfrich et al., 2014a) and TACS-MEG applications (Soekadar et al., 2013a; Neuling et al., 2015). For TMS-EEG, a recent study could convincingly show that provided appropriate artifact reduction procedures are followed, TMS-evoked potentials are absent from EEG in patients with extensive cortical lesions when damaged tissue is stimulated, but intact when the functional portion of cortex was targeted (Gossseries et al., 2015), suggesting that electrical artifacts can effectively be eliminated by existing procedures. Likewise, recent TACS-EEG studies evaluating event-related potentials in response to sensory stimuli (Helfrich et al., 2014b) or physiological brain activity patterns (in terms of topography and reactivity to eyes-open and closed conditions) (Neuling et al., 2015) suggest effective artifact reduction despite simultaneous electrical stimulation. In general, however, artifact reduction algorithms often require computationally heavy processing steps (including independent component analysis (ICA), source estimates or data interpolation). This allows for retrieving the EEG/MEG signal offline to the recordings but is yet incompatible with real-time analyses and interventions in many cases. For real-time applications, alternative neurophysiological read-outs or stimulation procedures circumventing the artifacts have therefore been used (see Section 6.1 below). Furthermore, while several procedures for artifact reduction exist, future research is needed for further evaluation and improvements.

In addition, control conditions should be designed to equate the potential sensory confounds of TMS and TACS, which may interact with brain oscillations by themselves (e.g., cause entrainment). Ideally, active controls should be used that are as similar as possible to the main condition in terms of the sensory component but less effective in regards to transcranial cortical stimulation. For TMS, this could consist of rotating coil orientations to a less effective direction of current flow (Thut et al., 2011) or the use of extracranial control sites like the shoulder blade (Herring et al., 2015) that provide comparable multisensory (auditory and tactile) inputs. For TACS, active control montages are likewise desirable, since the retina, inner ear, and peripheral sensory and vestibulo-cochlear nerves have low stimulation thresholds, as a result of which even subliminal (unperceivable) stimulation may affect brain oscillations and confound “transcranial” cortical effects (Utz et al., 2010; Schutter, 2016). Control montages can include extracerebral return electrodes, or when possible, the use of other stimulation frequencies that are behaviorally not relevant.

For the documentation of entrainment effects, control conditions as discussed above need to be implemented, because confounds during stimulation may also affect after-effects.

5. Open question: mechanisms of interventions

Fundamentally, the effect of rTMS and TACS on neuronal dynamics of oscillatory brain activity remains mostly unknown. This is because online monitoring of brain activity during stimulation is technically challenging and the extrapolations from the low number of studies using animal model preparations and computer simulations are not straightforward. Despite its limitation, the conceptual model which has gained the most traction for the response of brain networks to periodic (frequency-tuned) NTBS focuses on entrainment (Thut et al., 2011). Most generally spoken, entrainment refers to the behavior of an oscillating system to a periodic perturbation, where the system “locks” to the stimulation such that its frequency shifts to the frequency of the applied stimulation (or a harmonic/subharmonic). Certain conditions must be met for a periodic perturbation to accomplish successful entrainment. The stronger the intensity of stimulation, the broader is the range of frequencies (centered at the endogenous frequency in absence of stimulation) at which the network can be entrained. This principle is referred to as the Arnold tongues and has been well described for (quasi-)linear systems subjected to an external periodic force. It is important to note that while such Arnold tongue behavior has been shown in computer simulations (Ali et al., 2013; Herrmann et al., 2016), experimental evidence from animals or humans is limited due to the required number of stimulation intensities and frequencies that need to be evaluated.

For example for TMS, the relationship between stimulation intensity and potential phase resetting has not been sufficiently characterized yet. TMS at lower intensities or in the presence of strong endogenous oscillations may exert a phase-dependent effect while leaving the ongoing oscillation relatively unaffected. Conversely, at sufficiently high intensities or for weak endogenous oscillations TMS may phase-reset the circuits generating the endogenous oscillation and result in a TMS-locked oscillation. Importantly, in the latter case phase-reset would not necessarily be expected for the entire ongoing oscillation (observed from the summed potential/field EEG or MEG recordings), but rather for a circumscribed local population of stimulated neurons. While being largely unknown, the relationship between stimulation intensity and phase-reset/entrainment is of relevance for both the EEG-triggered informed open-loop and fully closed-loop approaches (see next section).

In addition, it is important to note that the underlying assumption of brain network oscillations to reflect measures generated by quasi-linear signals does not necessarily hold. For example, stimulation with periodic pulse trains at 10 Hz in epilepsy patients implanted with subdural electrode arrays for clinical monitoring revealed that entrainment may occur in the case of a relative weak endogenous oscillation, which is more susceptible to perturbations, whereas in states of pronounced endogenous oscillations the effect of the stimulation is not a shift to the stimulation frequency (as would be expected with entrainment), but rather an
increase in power at the endogenous frequency (Alagapan et al., 2016). This phenomenon can be easily explained by simple non-linear threshold models, but not by the more commonly used Arnold tongue framework (Alagapan et al., 2016). In conceptual agreement, in a reduced slice preparation that combined optogenetic activation of the network with electric field application, only weak optogenetic activation did the electric field enhance the activity at the stimulation frequency (Schmidt et al., 2014).

As to the mechanisms underlying changes in brain oscillation by amplitude-modulated TACS (as opposed to mono-sinusoidal TACS) in which the amplitude of a high-frequency carrier signal is modulated at a frequency of interest (Witkowski et al., 2016), it was discussed that the entrainment effect of such protocols might be related to non-linear properties of cell membranes (Goldman, 1943) resulting in rectification of the TACS signal. In this context, configurational changes of membrane proteins that lead to modifications of ionic binding sites and membrane permeability may play an important role.

6. Future perspective: optimization of the approach

While there is emerging evidence for the interest of informing NTBS by EEG/MEG in terms of timing and frequency, many of the reported effects still await replication and need to be evaluated as to whether their effects sizes lend themselves to clinical applications and/or can be further amplified. In this endeavor, the choice of optimal stimulation parameters will be important. While timing and frequency is informed by EEG/MEG, many other stimulation parameters are normally chosen arbitrarily (due to a lack of knowledge on how to guide them). Since the parameter space to choose from is almost infinite, including (i) intensity of stimulation for TMS and TACS, (ii) pulse form and coil orientation for TMS, (iii) electrode montage and stimulation waveform (sinusoidal, saw-tooth, rhythmic squared, i.e. pulsed, amplitude-modulation) for TACS, (iv) number of pulses or duration of stimulation, (v) stimulation of resting state vs. stimulation during a task etc., an exploration of the whole parameter space for finding the best parameter configuration is likely unfeasible. The development of optimization strategies will therefore be an important line of future research. For interactions with brain oscillations, two such strategies have been proposed, namely real-time closed-loop interventions and guidance by biological plausible models, as outlined below.

6.1. Data-driven optimization: real-time open-loop versus closed-loop approaches

Recent technical advances enable the use of informed open-loop and even fully closed-loop approaches, which evaluate the EEG (or in principle also MEG) signal in real-time to control the concurrent NTBS application accordingly (Bergmann et al., 2016; Zrenner et al., 2016): informed open-loop exploits amplitude and phase information of a specific ongoing oscillation in order to trigger stimulation in a temporally specific manner, however, without aiming to change the underlying oscillation. In contrast, a fully closed-loop approach aims to alter the targeted neuronal activity (Karabanov et al., 2016), e.g., by increasing or decreasing the amplitude of oscillatory brain activity or phase-locking it to the stimulation. See Fig. 5 for a schematic representation. Provided that brain oscillations are effective targets for NTBS, these approaches are expected to help optimizing interventions.

The first study using EEG-triggered TMS in an informed open-loop manner quantified motor cortical excitability and TEP changes during different phase-angles of the slow <1 Hz NREM sleep oscillation (Bergmann et al., 2012a). Nowadays, oscillatory phase angles can be assessed with even higher temporal precision and shorter time delays, thus also allowing to target faster oscillations in real-time (Zrenner et al., 2010, 2015, 2016; Triesch et al., 2015). It needs to be noted however that depending on sampling rates, communication protocols or data pre-processing steps (e.g. head localization in MEG), timing delays and jitter may occur, which can impede precise timing of stimulation.

To date, no closed-loop NTBS has been demonstrated using EEG that relies on simultaneous read-out and targeting of activity in neural population (Karabanov et al., 2016). For a recent study that relies entirely on activity in neural population but using read-out in stimulation-free intervals only see Lustenberger et al. (2016). This limitation is predominantly due to the considerable stimulation artifact in EEG recordings caused by either TMS or TACS, which interferes with real-time assessment of the oscillatory target parameters, once stimulation has started (for a detailed discussion see Bergmann et al., 2016). Instead, closed-loop strategies with EEG recordings have thus far employed either indirect read-outs of neural activity (e.g. concentrating on the behavioral consequences of the stimulation), or non-electrical forms of stimulation that circumvent stimulation artifact entirely. For instance, Ngo et al. (2013, 2015) delivered auditory tones for interventions during sleep. The tones were phase-triggered to slow sleep oscillations
with the aim to enhance these oscillations, facilitating overnight memory consolidation. In another example using motor cortex TACS to reduce tremor in Parkinson’s disease, actigraphy from the tremulous limb was adopted as a proxy for central neural activity (Brittain et al., 2013). The approach involved an initial open-loop followed by a closed-loop intervention that afforded several key advantages: First, knowledge of phase information for both TACS waveform and target oscillations (made possible by the proxy measure of central activity) permitted a direct measure of entrainment (Mehta et al., 2014, 2015; Brittain et al., 2015). Second, the phase-precession associated with open-loop stimulation permitted the construction of a stimulus response profile, in order to identify the most effective stimulation parameters for tremor suppression. Third, the phase of the target oscillation could then be used to directly inform real-time closed-loop (phase-locked) stimulus delivery. Indeed, stimulus response profiles revealed that (for tremor at least) it was the phase-difference between stimulation and tremor oscillations – rather than the phase of stimulation itself – which was the crucial factor in selectively suppressing or exacerbating the peripheral tremor. Of course, this approach relied on the hypothesis of a strong (causal) relationship between central oscillatory rhythms and peripheral outflow, surmised from prior functional connectivity studies (Timmermann et al., 2003). Yet despite this caveat, the closed-loop TACS saw a marked increase in effect-size relative to open-loop stimulation (Brittain et al., 2013). Finally, as an alternative real-time intervention approach, temporal interleaving of stimulation and recording epochs may also be considered. This approach has been shown to successfully control alpha oscillations using TACS in an early report (Boyle and Fröhlich, 2013) and was recently used to detect oscillatory transients in real-time (namely sleep spindle activity during NREM sleep) to then engage the target oscillation by individualized TACS limited to brief time epochs, enhancing not only sleep spindle activity but also overnight memory consolidation (Lustenberger et al., 2016). In addition, feasibility of NTBS during online MEG (instead of EEG) was successfully demonstrated and allowed a chronic stroke patient to modulate ipsilesional sensori-motor rhythms (SMR) while tDCS was applied to the ipsilesional sensori-motor cortex (Soekadar et al., 2013b). This study suggests that also MEG source activity-informed TACS will be feasible (requiring a real-time MEG system, though).

As a cautionary note, it needs to be mentioned that it is unknown how the sustained cumulative effects of closed-loop stimulation delivered under steady-state conditions (such as during bouts of tremor) would translate to behavioral paradigms, where fluctuating neural dynamics are constantly being reset and updated in a context and state-dependent manner. In addition, in several studies entrainment has been reported to be weak (Mehta et al., 2014, 2015; Brittain et al., 2015). Since open-loop protocols rely on assumptions about underlying mechanisms (e.g. steady entrainment of ongoing oscillations), there is a danger that such protocols will be undermined by unobserved temporal relationships and dynamic changes in the course of stimulation (e.g. due to homeostatic plasticity), even when TACS is delivered at the (subject-specific) natural resonance frequency of the targeted neuronal circuit. The advent of informed open-loop and closed-loop approaches linked to real-time electrophysiological readouts (such as EEG and/or MEG) therefore appears to offer a crucial step forward in optimizing stimulation protocols.

6.2. Model-driven optimization: prediction of effects with biologically plausible models and simulations

Another promising, emerging means for guiding intervention is computational modeling of stimulation effects, also referred to as computational neurostimulation (Bestmann, 2015) (for TMS see Rusu et al., 2014; Hartwigen et al., 2015; Triesch et al., 2015; for TACS see Fröhlich, 2015). These models implement a physiological component to work towards a mechanistic understanding of the NTBS-brain interaction, and are complementary to e.g. anatomically realistic finite element models that estimate actual intracranial field distributions (Opitz et al., 2015), which may be used to correct for substantial inter-individual variability in this measure (Opitz et al., 2016). In the modeling approach, mathematical models of the targeted neuronal networks can be subjected to stimulation in computer simulations allowing to take into account several parameters (e.g. spatial and temporal) of NTBS. In contrast to human or even animal studies, computational modeling therefore allows for the rapid evaluation of a large number of stimulation parameter combinations in terms of their effects on network dynamics. Such comprehensive parameterization has enabled the identification of entrainment of brain oscillations (due to the presence of Arnold tongue behavior, see Section 5 above), as a fundamental mechanism of target engagement by TACS (Ali et al., 2013; Herrmann et al., 2016). This is particularly important for TACS, where the small magnitude of the electric field delivered to the brain requires a synergistic interaction with endogenous network dynamics for the stimulation to have an effect (Fröhlich, 2015). In addition, mathematical models of sufficient biological plausibility enable the identification of cellular mechanisms that contribute to the effect of TACS. While there is certainly a broad range of computational models that can help to elucidate the interaction between the network dynamics targeted by stimulation and the applied stimulation, models that accurately model the non-linear dynamics for membrane voltage values around the action potential “threshold” may be of particular importance since this non-linearity likely enables the small changes in membrane voltage induced by NTBS to have an effect on the spiking of individual neurons and thus the network (Bonaiuto and Bestmann, 2015).

7. Future perspective: the promise of multimodal neuroimaging (TMS-EEG-fMRI)

As described above, electrophysiological methods such as EEG/MEG are capable of noninvasively measuring the temporal aspects of ongoing neuronal activity and their fluctuations, thereby capturing various parameters of neural oscillations, which may then be used to guide NTBS for optimizing when and how to stimulate. Complementing this focus on the temporal domain, functional magnetic resonance imaging (fMRI) – the established noninvasive method for measuring the spatial aspects of function-related neural activity – allows examining activity changes in localized brain regions and networks across the whole brain. Much has been written about the various neuroimaging tools and their complementary benefits, and it is widely accepted that combining them provides, at least conceptually, a rich and relatively complete view on brain function at the macroscale.

‘Multimodal imaging’ generally involves a two-way combination of tools, such as fMRI-EEG, fMRI-TMS, fMRI-TACS, EEG-TMS, or EEG/MEG-TACS, which can encompass either offline or online (i.e. simultaneous) combinations (Siblner et al., 2009; Reithler et al., 2011; Ziemann, 2011; Bergmann et al., 2016). Such simultaneous combinations are challenging, both technically and in terms of analysis, but have added value over and above offline combinations (Siblner et al., 2009). The reason for this is straightforward: in simultaneous setups, the same participant can be measured in the same environment, position and mindset with the same fluctuations of attention, comfort, and equivalent influence of other extraneous variables. This increases validity and eliminates certain sources of noise. More importantly, simultaneous combinations
allow us to uniquely address particular questions, and to approach data analysis in specific ways, in order to deliver more fundamental insights on the interaction of NTBS with brain activity and related behavior.

Here, we focus on the value of adding fMRI to EEG-TMS or EEG-TACS research (i.e. the three-way combination approach). In a feasibility study, Peters et al. (2013) combined commercially available hardware for simultaneous TMS-fMRI and simultaneous EEG-fMRI to evaluate safety/comfort and signal quality using a variety of tests on phantom and human volunteers. The three-way simultaneous combination of fMRI, EEG and TMS was shown to be safe, tolerable, and to provide good-quality signals. This grants new opportunities for future research to learn about how oscillatory activity in different frequency bands relates to activity and interactions in large-scale functional brain networks, and how this may be modulated by TMS/TCS. In a first approach to combine EEG, TACS and fMRI, Vosskuhl et al. (2016) showed that tACS at the individual alpha frequency – as determined by EEG – down-regulates the visually event-related BOLD response but not the ongoing BOLD activity.

Simultaneous fMRI-EEG already allows meaningful multimodal integration, and to correlate fluctuations in oscillatory parameters (e.g. increases/decreases of power in a particular frequency-band) to fluctuations in network BOLD signals (Debener et al., 2006; Scheeringa et al., 2009, 2011; Bergmann et al., 2012b). But because the temporal scales of EEG and fMRI are so far apart (operating in the range of milliseconds versus seconds, respectively), this approach has limitations. The simultaneous three-way combination of TMS-EEG-fMRI may offer a unique way to look deeper into the relation between oscillatory parameters such as phase, power, and coherence, and network activity as measured by fMRI. This setup allows the integration of the spatial and temporal domain, using one imaging method for indexing brain state (e.g. EEG to determine ongoing oscillation parameters such as power and phase), while the complementary imaging method can serve as the actual read-out measurement (e.g. TMS-induced BOLD fMRI network effects). By this means, the modulation of TMS-probed cortico-subcortical network effects (revealed by TMS-fMRI) can be assessed as a function of ongoing cortical oscillation parameters (as indexed by concurrent EEG). In other words, TMS pulses are used as system probes, inducing a network response measured by fMRI, depending on the EEG-measured oscillatory brain state at the time of the pulse. Besides providing fundamental insight on network functions (in particular if combined with behavioral assays), the three way approach should help the understanding of NTBS-effects (and their state-dependency) in important ways. More specifically, one can ask the question to what extent TMS-elicted BOLD responses throughout motor, perceptual, or cognitive networks are scaling with the momentary power, phase, or coherence of oscillations in functionally relevant frequency bands. Additionally, one can clarify how such EEG-fMRI network responses depend on functional state, such as rest versus task, different task conditions, or parametrically varied task loads, in order to provide fundamental insights into the relationship between oscillations, behaviorally relevant brain networks and NTBS effectivity.

8. Conclusion

We reviewed emerging ideas on how to work towards enhancing the specificity and effectivity of established NTBS protocols, based on the combination of NTBS mainly with EEG/MEG (and also fMRI). These ideas emphasize brain oscillations as key players in a number of fundamental circuit motifs that influence brain functions (Singer, 2009) and, as a consequence, constitute interesting targets for interventions. We identified three approaches that have been successfully used to interact with oscillatory brain activity: (1) triggering NTBS events to instantaneous phase- or power-values of ongoing EEG/MEG that reflect states of heightened excitability, which is promising for enhancing NTBS effectivity; (2) tuning NTBS to the known frequencies of specific task-relevant brain oscillations, in order to entrain these oscillations and promote the functions of the associated network; and (3) phase-triggering NTBS events to NTBS-entrained oscillatory brain activity, in order to potentiate approaches 1&2 by their combination. Initial results are promising, but further research is needed to document in more detail the electrophysiological underpinning of NTBS-induced network changes when guided by EEG/MEG for working towards a mechanistic account. The approach also affords the opportunity for implementing in future research both data- and model-driven optimization strategies (via real-time interventions and computational neurostimulation), which will be crucial for developing NTBS into an effective tool for experimental and clinical interventions into brain network activity and its (dys)functions. In brief, the outlined approach and ideas offer a framework for a hypothesis-driven, principled way of tailoring brain stimulation to interact with brain activity for shaping the brain-behavior relationship, constituting a promising new departure from conventional NTBS studies.

Acknowledgments

GT is supported by the Wellcome Trust [grant number 098434]. TOB is supported by the German Research Foundation via TR-SFB 654 (“Plasticity and Sleep”), and by the Hertie Foundation via the Hertie Institute for Clinical Brain Research. FF is supported by the National Institute of Mental Health of the National Institutes of Health under Award Number R01MH101547. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. AV-C is supported by the IHU-ICM-Translational initiative & Agence National de la Recherche Scientifique (ANR Génerique 2015 “Oscilloscopus”). ATS is supported by a grant from the Netherlands Organisation for Scientific Research (NWO VICI, grant number: 453-15-008). JSB is supported by the Medical Research Council (MR/N003446/1). HRS is supported by the Novo Nordisk Foundation (Interdisciplinary Synergy Programme Grant “BASICS” NNFI400C0011413). CSH and UZ are supported by the German Research Foundation (HE 3353-8/2 and ZI 542/7-1, respectively).

Conflict of interest: GT has received honoraria as editor from Wiley Publishers. FF is the lead inventor of IP filed by UNC. The clinical studies performed in the Frohlich Lab have received a designation as conflict of interest with administrative considerations. FF is the founder, CSO, and majority owner of Pulvinar Neuro LLC. HRS has served on a scientific advisory board for Lundbeck A/S, Valby Denmark, and has received honoraria as speaker from Biogen Idec, Denmark A/S, Genzyme, Denmark and MerckSerono, Denmark, has received honoraria as editor from Elsevier Publishers, Amsterdam, The Netherlands and Springer Publishing, Stuttgart, Germany, has received travel support from MagVenture, Denmark, and has received a research grant from Biogen-idec. UZ has received personal fees from Biogen Idec GmbH, Bayer Vital GmbH, Bristol Myers Squibb GmbH, CorTec GmbH, Medtronic GmbH, and grants from Biogen Idec GmbH, Servier, and Janssen Pharmaceuticals NV, outside of the submitted work. CSH has received honoraria as editor from Elsevier Publishers, Amsterdam. The remaining authors have no conflicts of interest.
Dugue L, Marque P, Vanrullen R. The phase of ongoing oscillations mediates the


