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Evidence-based guidelines on the therapeutic use of transcranial direct current stimulation (tDCS)

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List of abbreviations
AD: Alzheimer’s disease; ARAT: action research arm test; BDI: Beck depression inventory; BI: Barthel index; DC: direct current; DLPFC: dorsolateral prefrontal cortex; EEG: electroencephalography; ERP: event-related potentials; FMA: Fugl-Meyer assessment; fMRI: functional magnetic resonance imaging; GABA: gamma-aminobutyric acid; HDRS: Hamilton depression rating scale; JTT: Jebsen-Taylor hand function test; M1: primary motor cortex; MADRS: Montgomery-Asberg depression rating scale; MCS: minimally conscious state; MEP: motor evoked potential; NMDA: N-methyl D-aspartate; OMCASS: Orgogozo MCA scale; PANSS: positive and negative symptoms scale; PD: Parkinson’s disease; PES: peripheral electrical stimulation; RCT: randomized controlled trial; rTMS: repetitive transcranial magnetic stimulation; SANS: Scale for the Assessment of Negative Symptoms; tACS: transcranial alternating current stimulation; tDCS: transcranial direct current stimulation; TMS: transcranial magnetic stimulation; tRNS: transcranial random noise stimulation; tsDSC: transcutaneous spinal direct current stimulation; UPDRS: unified Parkinson’s disease rating scale; V1: primary visual cortex; VAS: visual analogue scale; VRT: virtual reality training; VS: vegetative state.
Abstract
A group of European experts was commissioned by the European Chapter of the International Federation of Clinical Neurophysiology to gather knowledge about the state of the art of the therapeutic use of transcranial direct current stimulation (tDCS) from studies published up until September 2016, regarding pain, Parkinson's disease, other movement disorders, motor stroke, poststroke aphasia, multiple sclerosis, epilepsy, consciousness disorders, Alzheimer’s disease, tinnitus, depression, schizophrenia, and craving/addiction. The evidence-based analysis included only studies based on repeated tDCS sessions with sham tDCS control procedure; 25 patients or more having received active treatment was required for Class I, while a lower number of 10-24 patients was accepted for Class II studies. Current evidence does not allow making any recommendation of Level A (definite efficacy) for any indication. Level B recommendation (probable efficacy) is proposed for: (i) anodal tDCS of the left primary motor cortex (M1) (with right orbitofrontal cathode) in fibromyalgia; (ii) anodal tDCS of the left dorsolateral prefrontal cortex (DLPFC) (with right orbitofrontal cathode) in major depressive episode without drug resistance; (iii) anodal tDCS of the right DLPFC (with left DLPFC cathode) in addiction/craving. Level C recommendation (possible efficacy) is proposed for anodal tDCS of the left M1 (or contralateral to pain side, with right orbitofrontal cathode) in chronic lower limb neuropathic pain secondary to spinal cord lesion. Conversely, Level B recommendation (probable inefficacy) is conferred on the absence of clinical effects of: (i) anodal tDCS of the left temporal cortex (with right orbitofrontal cathode) in tinnitus; (ii) anodal tDCS of the left DLPFC (with right orbitofrontal cathode) in drug-resistant major depressive episode. It remains to be clarified whether the probable or possible therapeutic effects of tDCS are clinically meaningful and how to optimally perform tDCS in a therapeutic setting. In addition, the easy management and low cost of tDCS devices allow at home use by the patient, but this might raise ethical and legal concerns with regard to potential misuse or overuse. We must be careful to avoid inappropriate applications of this technique by ensuring rigorous training of the professionals and education of the patients.

Keywords:
Cortex; indication; neurological disease; neuromodulation; noninvasive brain stimulation; psychiatric disease; tDCS; treatment.
Highlights

1. A group of European experts reviewed current evidence for therapeutic efficacy of tDCS.
2. Level B evidence (probable efficacy) was found for fibromyalgia, depression and craving.
3. The therapeutic relevance of tDCS needs to be further explored in these and other indications.
1. Principles and mechanisms of action of transcranial direct current stimulation

 Alterations of neuroplasticity and cortical excitability are important pathophysiological factors in many neuropsychiatric diseases. Thus, to modify cortical activities by using non-invasive brain stimulation (NIBS) might be a valuable therapeutic approach. One of these NIBS approaches is transcranial direct current stimulation (tDCS). Fifty years ago, it has been demonstrated in anesthetized rats that neural activity and cortical excitability could be modified by the application of direct current on the sensorimotor cortex, these effects depending on stimulation polarity and persisting for hours after the end of stimulation (Bindman et al., 1964). A few years later, it was established that a current flow sufficiently large to achieve physiological and functional effects could also be induced into the brain via transcranial application of such direct currents in both healthy subjects and patients suffering from psychiatric diseases (Rush and Driscoll, 1968; Dymond et al., 1975; Lolas, 1977). However, mainly due to the lack of relevant tools to assess its heterogeneous effects, this technique was nearly forgotten in the following years. About 15 years ago, tDCS was re-discovered as a tool to modulate human brain activity and its physiological effects started to be systematically explored (Priori et al., 1998; Nitsche and Paulus, 2000).

 The primary effect of tDCS on neurons is a subthreshold shift of resting membrane potentials towards depolarization or hyperpolarization, depending on current flow direction relative to axonal orientation (Bindman et al., 1962, 1964; Purpura and McMurtry, 1965; Gorman, 1966). Delivered to the motor cortex of healthy subjects, anodal tDCS increases the excitability of the underlying cortex, as revealed by an increase in the amplitude of motor evoked potential (MEP) to transcranial magnetic stimulation (TMS), whereas cathodal tDCS decreases it (Nitsche and Paulus, 2000). Stimulation of short duration (several seconds) is sufficient to induce these excitability changes, which however does not relevantly outlast the stimulation period (Priori et al., 1998; Nitsche and Paulus, 2000). A longer stimulation duration (several minutes) induces excitability changes that can last for one hour or more (Nitsche and Paulus, 2001; Nitsche et al., 2003b; Priori, 2003). As shown by a navigated TMS study, M1 excitability changes become steadily significant after the end of tDCS application rather than during stimulation (Santarnecchi et al., 2014). In these studies cited above, M1 was primarily targeted, but similar tDCS effects were obtained for the stimulation of visual (Antal et al., 2004) and somatosensory (Matsunaga et al., 2004) cortices.

 Overall, calcium-dependent synaptic plasticity of glutamatergic neurons is thought to play a key role in the outlasting neuroplastic mechanism of action of tDCS, since blockade of N-methyl D-aspartate (NMDA) receptors diminishes tDCS effects (Liebetanz et al., 2002; Nitsche et al., 2003a). In addition, tDCS can locally reduce gamma-aminobutyric acid (GABA) neurotransmission, regardless of stimulation polarity (Stagg et al., 2009) and this may also impact on glutamatergic plasticity due to the close relationship between the two neurotransmitters.
Beyond local effects, connectional effects of tDCS have also been described. Neuronal networks respond to DC fields even more sensitively than single neurons (Francis et al., 2003) and tDCS may interfere with functional connectivity, synchronization, and oscillatory activities in various cortical and subcortical networks. This has been shown for tDCS delivered to M1 (Polanía et al., 2011a,b, 2012), the prefrontal cortex (Keenser et al., 2011a), or during slow-wave sleep (Marshall et al., 2004).

In addition, since tDCS modulates resting membrane potential at the synaptic level but more generally along the whole axons, this may result in non-synaptic effects, which might also contribute to the long-lasting after-effects of tDCS (Ardolino et al., 2005). These non-synaptic mechanisms of tDCS might be based on changes of conformation and function of various axonal molecules, involved in transmembrane ion conductance, membrane structure, cytoskeleton, or axonal transport, when exposed to a DC field (Jefferys, 1995).

Another important consideration in understanding tDCS effects is that almost all tissues and cells are sensitive to electric fields and, therefore, tDCS might also elicit changes in non-neuronal tissues in the brain, including endothelial cells, lymphocytes, or glial cells (Ruohonen and Karhu, 2012). These non-neuronal effects, which have not been systematically explored to date, could also be involved in the therapeutic action of tDCS. In patients with cerebral diseases, besides neuronal damage, other important pathological processes may exist in the axonal microenvironment, such as inflammation. Beyond neuroinflammatory diseases, such as multiple sclerosis, inflammation in the central nervous system has been implicated in several neuropsychiatric conditions, such as in the progression of neurodegeneration in Alzheimer’s disease (AD) (Heneka et al., 2015). Hence, through its possible influence on the inflammatory response, tDCS might theoretically impact on the course of the disease. Also relevant in AD, one should note that the conformation of beta-amyloid and other pathological proteins can be changed when exposed to appropriate electric fields (Toschi et al. 2009), possibly modifying their susceptibility to degradation. In addition, DC fields can enhance axonal regeneration and neurite outgrowth (Fehlings and Tator, 1992; Wood and Willits, 2006, Pelletier et al., 2014) and therefore hypothetically improve functional recovery. Finally, tDCS could also be useful to limit the vicious circle of autodestructive events due to the increased Ca\(^{2+}\) influx resulting from excessive membrane depolarization or intra-axonal Na\(^+\) overload in the context of ischemia or energetic resource failure (Lefaucheur, 2009). In other words, tDCS might be able to influence several pathological processes and pathogenetic cascades in the central nervous system, well beyond the sole change of neuronal excitability.

However, the classically assumed effects of tDCS polarity on cortical excitability (i.e. excitatory anodal tDCS versus inhibitory cathodal tDCS), with primarily concerned motor cortex stimulation, cannot be considered as a general rule, since numerous factors can turn facilitatory changes into inhibitory, and vice-versa. The local cellular influence of DC fields into the brain is complex, depending on the distance and orientation of the axonal or somatodendritic axis with respect
to the electric field (Purpura and McMurtry, 1965; Gluckman et al., 1996; Bikson et al., 2004). Axon terminals are thought to be two-three times more susceptible than somas to tDCS-induced polarization, but radial or tangential current flow did not determine the same changes in synaptic efficacy at axon terminal level, according to stimulating electrode polarity (Rahman et al., 2013). More precisely, axonal orientation could determine whether the DC field is excitatory or inhibitory, whereas dendritic orientation could affect the magnitude but not the direction of DC resulting effects (Kabakov et al., 2012). In addition, modeling studies showed that a focal cathodal stimulation applied on the surface of a gyrus only or preferentially activates ‘horizontal’ fibers with a directional component parallel to the electrode surface, whereas anodal stimulation rather excites fibres in the underlying cortex that are perpendicular to the electrode surface (Manola et al., 2005; Holsheimer et al., 2007a,b). Finally, even if DC fields induce coherent depolarizing or hyperpolarizing effects on neuronal populations with respect to fiber orientation and electrode polarity, the resulting physiological effect of the stimulation depends on whether the affected network is dominantly inhibitory or excitatory.

Baseline activity of the neural networks and afferent synaptic inputs exposed to DC fields also have a relevant influence, according to the concept of metaplasticity (Abraham and Tate, 1997; Turrigiano and Nelson, 2004; Müller-Dahhaus and Ziemann, 2014), especially following the ‘Bienenstock–Cooper–Munro (BCM) model” (Bienenstock et al., 1982). This model assumes that synaptic depression is more likely occurring when postsynaptic activity is high, whereas synaptic potentiation is more likely occurring when postsynaptic activity is low. Thus, tDCS-induced plasticity differs whether tDCS is applied in a passive, relaxed subject or in a subject performing cognitive or motor task (Antal et al., 2007). Non-linear effects of tDCS have been described, depending on the relationship between tDCS parameters and the responsivity of various subtypes of neurotransmitter receptors and ion channels (Nitsche et al., 2012), which might entail a sudden lack of proportionality in stimulus-response relationships. In turn, neurotransmitters, especially dopamine, can exert a dosage-dependent, but non-linear effect on tDCS-induced plasticity (Monte-Silva et al., 2010b).

As underlying mechanism, it has been speculated that activation of neurons not only changes their membrane potential and firing rate but also reduces membrane resistance. This loss of resistance, or increase of conductance, may have a key importance for tDCS effects since electric fields can cause larger changes in transmembrane voltage in resting neurones with low membrane conductance than in active neurones with high membrane conductance (Paulus and Rothwell, 2016). Therefore, tDCS effects depend on complex spatial relationships between the stimulated active target field, its projection areas, the resting surrounding structures, the pathological alterations of transmitter systems, the medication taken by the patients (Brunoni et al., 2013a), and individual genetic polymorphisms (Plewnia et al., 2013).

One should also consider the major influence of stimulation parameters, such as intensity (Batsikadze et al., 2013) and session repetition timing (Monte-Silva et al., 2010a, 2013), which reflect some aspects of the non-linear relationship between tDCS settings and the biological effects produced.
For example, increasing the strength and duration of stimulation may enhance its efficacy in given applications (e.g., Shekhawat et al., 2016), but this cannot be considered as a general rule. In fact, increasing tDCS intensity might also shift the direction of excitability changes, e.g., doubling intensity from 1 mA to 2 mA can switch inhibition produced by cathodal tDCS of M1 into excitation (Batsikadze et al., 2013). In addition, the induced electric field spreads and goes deeper into the brain as intensity increases, able to modify the nature of the recruited neural networks and therefore the resulting biological and clinical effects unexpectedly (Lefaucheur, 2012). Finally, it is crucial to take into account that even little variations of electrode size area, shape, or placement (montage) can strongly influence the “diffusion” of the current and the geometry of the induced DC fields into the brain (Miranda et al., 2009; Faria et al., 2011; Sadleir et al., 2012; Saturnino et al., 2015). Therefore, the clinical changes provided by tDCS may strongly vary according to many technical and neuroanatomical considerations and relationships (Priori, 2003; Datta et al., 2012; Parazzini et al., 2012; Brunoni et al., 2014c; Ho et al., 2014; Galletta et al., 2015; Senço et al., 2015). All these factors could explain the variable results provided by tDCS trials in pathological conditions. They should be taken into account for designing stimulation protocols and interpreting the effects of tDCS when applied in patients with neuropsychiatric diseases. Personalized therapeutic application of brain stimulation techniques might be developed in the future, especially in the light of new findings, which show profound interindividual variability of cortical excitability changes in response to cortical stimulation using tDCS (Lopez-Alonso et al., 2014, 2015; Wiethoff et al., 2014; Strube et al., 2015), as also shown for repetitive transcranial magnetic stimulation (rTMS) using either conventional protocols (Sommer et al., 2002) or theta burst paradigms (Hamada et al., 2013).

2. Clinical applications of tDCS: literature data analysis

For each potential clinical indication of tDCS, a bibliographic search was carried out by several experts independently, using specific keywords that will be specified at the beginning of each section. Each expert then proceeded to a critical reading of all selected publications in order to classify them according to the following criteria, derived from those proposed by the European Federation of Neurological Societies (Brainin et al., 2004). First, the studies were classified (I to IV) according to decreasing value of evidence. A **Class I study** is an adequately data-supported, prospective, randomized, placebo-controlled clinical trial with masked outcome assessment in a representative population (n≥25 patients receiving active treatment). It should include (a) randomization concealment; (b) clearly defined primary outcomes; (c) clearly defined exclusion/inclusion criteria; (d) adequate accounting for dropouts and crossovers with numbers sufficiently low to have minimal potential for bias, and (e) relevant baseline characteristics substantially equivalent among treatment groups or appropriate statistical adjustment for differences. A **Class II study** is a randomized, placebo-controlled trial performed with a smaller sample size (n≥10, but <25) or that lacks at least one
of the above-listed criteria a–e. **Class III studies** include all other controlled trials, with less than 10 patients, or more than 10 patients, but having methodological limitations in the experimental design, control procedures, or objectives. **Class IV studies** are uncontrolled studies, case series, and case reports.

With the aim of establishing evidence of efficacy of a given tDCS protocol in any specific therapeutic indication, the experts then compared their respective classifications until they reached a consensus and applied these to the levels of evidence A to C, as follows, without considering Class IV studies. **Level A** ("definitely effective or ineffective") requires at least two convincing Class I studies or one convincing Class I study and at least two convincing Class II studies. **Level B** ("probably effective or ineffective") requires one Class I study and less that two Class II studies, or at least two convincing Class II studies, or one convincing Class II study and at least two convincing Class III studies. **Level C** ("possibly effective or ineffective") requires one convincing Class II study and less that two Class III studies or at least two convincing Class III studies. No recommendation is made on one isolated Class I, II, or III study.

For this study, only placebo-controlled studies (using sham tDCS, according to either crossover or parallel-arm design), based on repeated daily sessions and including at least 10 patients receiving an "active" tDCS protocol, were retained for further analysis. The restriction to repeated session protocols was justified by the fact that the present work addresses the therapeutic effect of tDCS in chronic disorders, which requires tDCS sessions to be repeated to produce sustained effects.

Regarding grading of the "level of evidence", we always considered recommendation for a specific indication (with the patients having the same disease with the same type of clinical features), using a specific stimulation protocol (with the same anode/cathode montage, irrespective of stimulation intensity (1-2 mA) or duration and number of sessions). In addition, single-center studies with the same indication and methodology coming from a given research group were only considered once (according to their best class).

For each indication, only clinical results reported in controlled studies, published before the end of the bibliographic search (September 1st, 2016), based on repeated tDCS sessions with sham tDCS control procedure, and including at least 10 patients receiving active stimulation, were included in the evidence-based analysis. If at least two comparable studies with similar clinical indication and tDCS montage were published by independent research groups, data were summarized in a Table.

These Tables give the number of patients who actually received tDCS therapy, excluding dropouts. In trials with parallel arms, the respective number of patients in the active and control groups are indicated. In the “Results” column, the main results are usually summarized as a function of the significance of the effect of active tDCS versus control condition. Following this analysis, we propose an overview of the level of evidence that can be currently recommended for the therapeutic application of tDCS in a given clinical indication, according to specified parameters of stimulation, especially regarding electrode locations and montage.
In addition, to clarify certain points (e.g., possible mechanisms of action), to give a larger overview of the “therapeutic” potential of the technique, or to indicate the tracks of future developments, we were led to quote in the text and discuss various studies, despite their small sample size or their open-label or single-session design. These studies, however, were not taken into account in the evidence-based analysis.

3. Pain

The literature review included studies related to ongoing chronic pain, or acute postoperative pain, and therefore excludes publications on the use of tDCS to relieve pain experimentally induced in healthy subjects, reviewed elsewhere (Mylius et al., 2012). A PubMed search [keywords: tDCS AND (pain OR migraine)] identified 269 papers, including 62 original clinical studies covering 1,426 patients. In all indications, we first screened literature data for original clinical trials, excluding any other papers, such as reviews, editorials, or experimental studies regarding animals or healthy subjects. From these 62 clinical trials, 17 included less than 10 patients. The types of pain syndrome were as follows:

- central neuropathic pain (144 patients), including central poststroke pain (2 studies) and medical or traumatic spinal cord injury (7 studies);
- peripheral neuropathic pain (95 patients), including peripheral neuropathy (1 study) and radiculopathy (1 study);
- musculoskeletal pain (excluding low back pain) (316 patients), including fibromyalgia (11 studies), myofascial pain (3 studies), and plantar fasciitis (2 studies);
- migraine (265 patients, 8 studies);
- orofacial pain (68 patients), including trigeminal neuralgia (2 studies), temporomandibular disorder (2 studies), and various orofacial pain syndromes (1 study);
- low back pain (266 patients, 5 studies);
- abdominal or pelvic pain (40 patients), including inflammatory bowel syndrome (1 study), endometriosis (1 study), and various types of pelvic pain (3 studies);
- postoperative pain (147 patients, 4 studies);
- miscellaneous syndromes (85 patients), including phantom limb pain (3 studies), multiple sclerosis (1 study), arthralgia (1 study), drug toxicity (1 study), and mixed etiologies (2 studies).

Generally, the tDCS target was M1, the left dorsolateral prefrontal cortex (DLPFC), or the primary visual cortex (V1) for migraine. In most studies, anodal stimulation was applied to M1 of the hemisphere contralateral to pain (in case of focal or lateralized pain) or the dominant (left) hemisphere (in case of more diffuse pain). M1 was usually defined as the location of the C3/C4 electrode in the International 10-20 system for EEG electrode placement. In the largest (but negative) tDCS study to date in the pain domain, concerning 135 patients with low back pain (Luedtke et al., 2015), the M1 target was rather defined as the motor hotspot of the hand determined by recording abductor digiti
minimi MEPs to single-pulse TMS. In almost all studies with the anode intended to be placed over M1 or the left DLPFC, the cathode was positioned over the contralateral supraorbital region. Conversely, in at least five studies, both M1 and DLPFC were stimulated concomitantly or subsequently, or the cathode was positioned in place of the anode over the intended cortical target, including cathodal stimulation of V1 to treat migraine (Antal et al., 2011).

It has been suggested that M1 anodal stimulation may reduce pain by activating various neural circuits present in the precentral gyrus, which would be afferents or efferents that connect structures involved in sensory or emotional component of pain processing, such as the thalamus or the DLPFC, or by facilitating descending pain inhibitory controls (Lefaucheur, 2006; Nguyen et al., 2011). In migraine, cathodal stimulation of V1 is thought to decrease its oversensitivity or increased responsiveness at the origin of headache.

In most studies, stimulation was delivered at an intensity of 2mA with an electrode size of 35cm². The duration of the daily stimulation session ranged from 10 min to 20 min, repeated for up to 20 sessions. The effect of a single tDCS session was also assessed in several studies, showing significant pain relief, except in two studies (Luedtke et al., 2012; Dubois et al., 2013). In these latter studies, both anodal and cathodal tDCS of M1 had no impact on acute pain provoked by noxious thermal and electrical stimuli superimposed to chronic low back pain (Luedtke et al., 2012) or postoperative pain in lumbar spine surgery (Dubois et al., 2013). The most commonly used protocol consisted of 20-min anodal stimulation of M1 for five consecutive days, which may lead to significant analgesic after-effects lasting for 2 to 6 weeks (Fregni et al., 2006e; Valle et al., 2009; Antal et al., 2010; Kim et al., 2013).

A responder is usually defined as a patient experiencing pain relief by more than 30-50% on a visual analogue scale (VAS) (Klein et al., 2015). It is still unclear whether tDCS treatment produces different levels of relief in different types of pain. Analgesic effects of anodal tDCS of M1 have been reported in various neuropathic pain syndromes of either central or peripheral origin, but especially located at the lower limbs (Table 1). Regarding this pain location, analgesic effects of tDCS have been reported in one class II study (Kim et al., 2013) and three class III studies (studies with active group of at least 10 patients, but sham group of less than 10 patients or combining tDCS with another intervention) (Fregni et al., 2006a; Soler et al., 2010; Yoon et al., 2014), but two class II studies did not show any significant difference in pain intensity reduction between active and sham condition (Wrigley et al., 2013; Attal et al., 2016). Finally, one class II study was ambiguous, reporting similar global pain relief after active and sham tDCS, but more responders after active vs. sham tDCS (Souto et al., 2014) (8/10 vs. 3/10). Therefore, we can only propose a recommendation of Level C (possible efficacy) regarding the analgesic effect of anodal tDCS of M1 applied contralaterally to the pain side or on the left hemisphere in patients with neuropathic pain at the lower limbs, at least when pain is secondary to spinal cord lesion. We can further recommend performing tDCS sessions of 20-min duration for at least 5 consecutive days, using 2 mA intensity and 35 cm² electrode size.
In fibromyalgia, a number of publications reported positive results in favor of anodal tDCS of the left M1 or DLPFC, but all coming from a single team, regarding the application of either a classical “bipolar” montage (Fregni et al., 2006e; Roizenblatt et al., 2007; Valle et al., 2009; Riberto et al., 2011) or a presumably more focal, multipolar Laplacian montage (Mendonca et al., 2011; Villamar et al., 2013; Castillo-Saavedra et al., 2016). More recently, three additional class II studies were published by independent teams, showing a greater pain relief produced by anodal stimulation of the left M1 compared to sham tDCS (Fagerlund et al., 2015; Foerster et al., 2015; Jales Junior et al., 2015). Overall, these studies reported a mean reduction of pain intensity ranging between 14 and 58% compared to baseline at the end of the tDCS protocol (5 to 10 sessions), which remained statistically significant up to one or two months after the end of the intervention. Short-lasting effects that were not prolonged for more than one week after 5 days of 1 mA anodal tDCS were observed in patients with focal myofascial pain in the shoulder (Sakrajai et al., 2014). Pain relief was associated with an improvement in the quality of life in most tDCS studies of patients with fibromyalgia syndrome. Therefore, a recommendation of Level B (probable efficacy) can be proposed regarding the analgesic effect of anodal tDCS of the left M1 in fibromyalgia. However, the clinical relevance of the intensity of pain relief induced by anodal tDCS of the left M1 in this condition is clearly debated (Fagerlund et al., 2015). A recent open-label trial showed a clinically significant benefit (50% pain reduction associated with quality of life improvement) in 7/14 patients treated for 6 weeks by tDCS delivered over M1 using a multipolar Laplacian montage (Castillo-Saavedra et al., 2016), but these results remain to be replicated by other teams independently. It is also important to determine whether the analgesic effects of tDCS in fibromyalgia syndrome can be distinguished from potential concomitant effects on depression and anxiety. Finally, we have no practical guidelines on how to design a maintenance protocol for providing long-term therapeutic effects of tDCS in this chronic disease.

In other pain conditions, there are no replicated convincing studies showing beneficial effects of tDCS applied to M1 or the left DLPFC provided by independent teams and therefore, no recommendation can be made. For example, there are only single positive studies regarding the value of repeated sessions of anodal tDCS of M1 contralateral to pain in trigeminal neuralgia (Hagenacker et al., 2014), pain related to multiple sclerosis (Mori et al., 2010), or irritable bowel syndrome (Volz et al., 2016). In pain related to temporomandibular disorder, one study using multipolar montage (2 cathodes and 2 anodes) reported beneficial effects (Donnell et al., 2015), whereas no difference was found in pain relief between active and sham condition in another study using a more classical bipolar tDCS montage with the anode over M1 (Oliveira et al., 2015). In chronic low back pain, the two published studies using repeated sessions of anodal tDCS of M1 were negative, the first one being based on a very small sample size (8 patients, therefore not retained in the analysis) (O’Connell et al., 2013), but the other one (Luedtke et al., 2015) including the largest series to date (135 patients) about tDCS studies in the pain domain.
With regard to other conditions, most studies had methodological limitations, such as small sample size or single-session design. For example, in a small group of patients, a single session of anodal tDCS of the left DLPFC performed just after endoscopic retrograde cholangiopancreatography was found to reduce pain intensity and morphine consumption after intervention (Borckardt et al., 2011). However, a subsequent study showed that a similar protocol (but with a different cathode placement) was unable to replicate these results in a larger series of patients undergoing gastric bypass surgery (Dubois et al., 2013). Later, in two studies, the group of Borckardt et al. reported the efficacy of a tDCS protocol consisting of four 20-min sessions performed from the immediate postoperative time to the day after surgery, with the anode over a region close to the leg area of the motor cortex (C1/C2 or Cz electrode site) and the cathode over the right DLPFC (F4 electrode site) in series of 39 patients undergoing total knee arthroplasty (Borckardt et al., 2013) and 27 patients undergoing lumbar spine procedure (Glaser et al., 2014). In both studies, tDCS was able to reduce postoperative opioid requirements. These results still remain to be replicated by another group.

In some indications, the value of the left DLPFC target was compared to that of M1 in terms of analgesic efficacy. The results were either similar (Valle et al., 2009) or in favour of M1, e.g., in patients with fibromyalgia (Fregni et al., 2006e) or diabetic polyneuropathy (Kim et al., 2013).

Finally, regarding migraine, the first positive results were reported by using cathodal tDCS of V1 (Antal et al., 2011; Wickmann et al., 2015). Other positive results were reported using anodal tDCS of M1 in two studies. One study included more than 10 patients receiving active tDCS and showed a reduction in pain intensity, number of attacks, and medication at the end of the intervention and up to 8 weeks after active tDCS (Auvichayapat et al., 2012). However, the other study was a preliminary report of results obtained in 8 patients, in which a reduction in the intensity of pain and the duration of migraine episodes was only observed in a delayed follow-up (Dasilva et al., 2012). Therefore, no recommendation can be made for this indication.

As a conclusion, the present level of evidence for the analgesic effect of M1 stimulation is weaker for tDCS than for rTMS (Lefaucheur et al., 2014). However, regarding this comparison, the most striking point is the fact that the best level of evidence for tDCS efficacy were found in different indications (fibromyalgia and neuropathic pain in the lower limbs) from that of M1 rTMS, which is known to be more prone to relieve chronic neuropathic pain at the face or upper limbs (Lefaucheur et al., 2004, 2006). One may speculate that tDCS and rTMS, although targeting the same motor cortical region, do not produce pain relief through the same mechanisms of action. This hypothesis is reinforced by the report of a patient with chronic refractory neuropathic pain who did not respond to high-frequency rTMS of M1, but was subsequently improved by repeated sessions of anodal tDCS of M1 in the long term (Hodaj et al., 2016). It has been suggested that rTMS might activate similar networks as epidural motor cortex stimulation (Lefaucheur et al., 2010), of which analgesic effect is obtained according to the placement of cathode(s) over the precentral gyrus (Holsheimer et al., 2007a,b), in contrast to tDCS for which the precentral electrode is classically an anode. In addition, the
intensity of rTMS and epidural stimulation is sufficient to elicit additional action potentials in the activated circuits, whereas tDCS is assumed to modulate spontaneous firing (Lefaucheur, 2012). Thus, different mechanisms of action might subserve different indications for high-frequency rTMS and anodal tDCS of M1 in the pain domain. Only one sham-controlled study directly compared the analgesic efficacy of these two approaches (Attal et al., 2016). This study included patients with neuropathic pain due to lumbosacral radiculopathy and showed that high-frequency rTMS, but not anodal tDCS of M1 contralateral to the painful area (three daily sessions) was superior to sham stimulation. This result is surprising, since, as mentioned above, relief of neuropathic pain in the lower limbs was more frequently reported after tDCS than after rTMS in the literature. Finally, as an alternative in the treatment of refractory chronic pain, tDCS offers various advantages compared to rTMS, e.g., a low-cost equipment that can be used at home, as shown in a recent case report of long-term treatment of myofascial pain using a tDCS protocol controlled by tele-medicine (Pérez-Borrego et al., 2014). Sham-controlled protocols of repeated tDCS sessions performed at home by the patients themselves are currently under investigation (O’Neill et al., 2015).

Please insert TABLE 1

4. Parkinson's disease

In advanced Parkinson’s disease (PD), the emergence of fluctuations, dyskinesias, difficulties with gait and postural control, cognitive impairment and non-motor symptoms refractory to conventional therapy poses therapeutic challenges. The success of deep brain stimulation (DBS) and advances in the understanding of the pathophysiology of PD have raised interest in NIBS techniques as alternative therapeutic tool. The rationale for the use of NIBS draws from the concept that reversing abnormalities in brain activity and physiology thought to cause the clinical deficits may restore normal functioning. Currently the best evidence in support of this concept comes from DBS, which improves motor deficits, and modulates brain activity and motor cortex physiology, though whether a causal interaction exists remains largely undetermined. At present, tDCS offers a safe, easy-applicable method of NIBS, and there are now several published tDCS trials aimed at improving various clinical aspects in the domain of PD (Benninger and Hallett, 2015).

Actually, a PubMed search (keywords: tDCS AND Parkinson's disease) identified 72 papers, including 15 original clinical studies and 225 patients. One paper was a single case report (Kaski et al., 2014a), whereas 10 to 25 patients were included in the other studies. The objectives of these studies performed in parkinsonian patients were various and can be summarized as follows:

- to study the effect of tDCS preconditioning (priming protocol, 1 mA for 10 min) on the ability of a subsequent low-frequency rTMS protocol (1 Hz, 900 pulses) delivered over M1 to improve
motor performance, motor control, or gain kinematics (Grüner et al., 2010; Eggers et al., 2012; Von Papen et al., 2014);

- to study the effect of anodal tDCS of the left DLPFC on working memory (Boggio et al., 2006) or verbal fluency (Pereira et al., 2013);
- to study the effect of anodal tDCS of the left or right DLFPC on executive or cognitive functions (Doruk et al. 2014; Manenti et al., 2016);
- to study the effect of anodal tDCS of the left and right DLFPC on walking abilities (Manenti et al., 2014);
- to study the effect of anodal tDCS over various motor regions on gait and motor performance (Verheyden et al., 2013; Kaski et al., 2014a,b; Valentino et al., 2014; Costa-Ribeiro et al., 2016; Ferrucci et al., 2016), optionally combined with prefrontal stimulation (Benninger et al., 2010).

**tDCS effects on motor symptoms.** In the literature, there are only three sham-controlled therapeutic tDCS trials based on repeated sessions of tDCS delivered over the motor cortex and including at least 10 PD patients in the active tDCS condition: two were parallel-designed and the other crossover-designed randomized controlled trials (RCTs) (Table 2).

In one parallel-designed RCT (Benninger et al., 2010), the safety and efficacy of anodal tDCS applied to the motor and prefrontal cortices in 8 sessions over 2.5 weeks were investigated. TDCS exerted significantly beneficial effects on gait (10 m walking) and bradykinesia (assessed by a timed testing of sequential hand and arm movements) for the follow-up period of 3 months. However, the observed effects were small and still need to be confirmed in a larger study. Conversely, changes in the motor score of the Unified Parkinson’s Disease Rating Scale (UPDRS), reaction time, physical and mental well-being, and self-assessed mobility did not differ between tDCS and sham intervention.

In a crossover RCT (Valentino et al., 2014), a protocol of 5 sessions of anodal tDCS of M1 was found to provide a beneficial effect on gait, freezing of gait, and motor performance in 10 parkinsonian patients. These effects lasted for the follow-up period of 1 month. The long-lasting persistence of effects in the both RCTs points out to plasticity phenomena. In the context of PD, it is important to note that dopamine, in a complex balance between D1 and D2 receptor mediation, exerts dose-dependent consolidation-enhancing effects on tDCS-induced changes of excitability and NMDA receptor-mediated neuroplasticity in the human motor cortex (Nitsche et al., 2006, 2009a; Fresnoza et al., 2014). In a recent study based on a small sample of 9 patients, anodal tDCS applied bilaterally over M1 for five consecutive days was found to improve levodopa-induced dyskinesias, but not other motor symptoms (Ferrucci et al., 2016). Anodal cerebellar tDCS provided the same results as M1 tDCS.

In addition, beyond the proper effects of tDCS on motor control and cortical excitability, there is increasing evidence pointing to the enhancement of motor learning and long-term retention as the rationale for combining tDCS protocol with rehabilitative intervention in PD patients (Benninger et al., 2010). This approach is supported by data obtained in healthy subjects (Nitsche et al., 2003c; Reis et al., 2009) and it was first applied to promote motor recovery in stroke patients (Hummel et al.,
In PD, this was the subject of two recent studies. They showed prolonged improvement in gait or motor abilities following anodal tDCS over the lower limb motor / supplementary motor area combined with cueing gait training (Costa-Ribeiro et al., 2016a,b) or anodal tDCS over the DLPFC contralateral to the most affected body side combined with physical therapy (Manenti et al., 2016).

Acute effects of single tDCS sessions have also been reported in PD patients. In a crossover study, a single session of anodal tDCS of left M1, but not of cathodal tDCS of left M1 or anodal tDCS of left DLPFC, was found to improve motor function (UPDRS score and simple reaction time) (Fregni et al., 2006d). In another crossover study, a single session of anodal tDCS of M1 of the dominant hemisphere reduced the time to walk 10 m, but had no effects on other gait measures (Verheyden et al., 2013). Gait velocity was also found to be improved after a single session of anodal tDCS of a region roughly corresponding to lower limb motor region with the cathode positioned at the inion (Kaski et al., 2014a,b).

tDCS effects on nonmotor symptoms. Cognitive impairment is prevalent in advanced PD and a major cause of disability and institutionalization, but fronto-executive dysfunction may manifest early. In a crossover study including 18 PD patients, a single session of anodal tDCS of left DLPFC, but not of left M1, improved performance in a working memory task, only at the stimulation intensity of 2 mA (Boggio et al., 2006). In another crossover study including 16 PD patients, a single session of anodal tDCS improved verbal fluency, associated with large-scale functional changes between connected brain areas, more significantly when the stimulation was applied to the left DLPFC than to the left temporo-parietal cortex (TPC) (Pereira et al., 2013). Finally, in a sham-controlled study of 20 PD patients with parallel-arm design, a protocol of 10 sessions of anodal tDCS delivered over the DLPFC contralateral to the most affected body side during a physical therapy program was found to increase cognitive performance and verbal fluency only in the active arm group (Manenti et al., 2016). The cognitive improvement was significantly lasting at 3-month follow-up, while an improvement in motor abilities and a reduction of depressive symptoms was similarly observed in both active and sham groups at the end of the treatment.

Conclusion. The currently published studies suggest a potential impact of anodal tDCS of M1 on gait and motor symptoms in PD patients, but do not provide sufficient evidence for a recommendation, because of heterogeneity in the targeted motor area or the outcome measures, for example. The clinical effects reported so far appeared to be negligible regarding functional independence and quality of life. Other results concerned small series of patients and mostly single-session protocols with short-lasting effects. Various montages and designed cortical targets remain to be investigated in larger studies. Combined approaches of tDCS with a rehabilitative intervention or priming strategies may potentiate the therapeutic efficacy (Benninger and Hallett, 2015). There is also a need to evaluate the value of tDCS protocols to act on nonmotor parkinsonian symptoms at a clinically meaningful level.
5. Other movement disorders

Actually, published tDCS studies on movement disorders other than PD are rare and have been recently reviewed (Ferrucci et al., 2016). There are only two sham-controlled crossover studies using repeated tDCS sessions. The first study concerned 8 patients with essential tremor who completed a tDCS protocol for 10 days with two cathodes placed over both cerebellar hemispheres and two anodes positioned over both prefrontal areas (Gironell et al., 2014). In this study, any short- or long-lasting clinical benefits in any outcome measure was observed after active stimulation compared to the sham condition. The second study was a case report of 2 patients with Tourette's syndrome who benefitted from a 5-day protocol of cathodal tDCS of left M1 (with the anode over the right deltoid muscle) (Mrakic-Sposta et al., 2008). Active stimulation significantly decreased the number of motor and phonic tics in both patients at the end of the 5-day treatment.

It is for dystonia that there would be more relevance to find tDCS studies. Indeed, dystonia is a heterogeneous disorder characterized by impaired motor control related to a loss of inhibition, sensory dysfunction, and alterations of synaptic plasticity leading to abnormal sensorimotor integration (Quartarone and Hallett, 2013). Clinical evidence support efficacy of DBS, whereas beneficial effects of NIBS, if any, are modest and short-lasting (Benninger et al., 2015; Cho and Hallett, 2016). Regarding tDCS and dystonia, a PubMed search (keywords: tDCS AND dystonia) identified 32 papers, including 15 original clinical studies and 147 patients. Thus, the amount of published data is not far from that of PD. However, most studies are case reports or small case series. Approximately only half of the studies included at least 10 patients. In addition, the studies were often issued from the same group, for specific clinical conditions, which were:

- musician's dystonia (77 patients; 7 studies);
- writer's cramp (35 patients; 4 studies);
- cervical dystonia (2 patients; 2 studies);
- pediatric cases of primary and secondary generalized dystonia or hemidystonia, including secondary to cerebral palsy (33 patients; 3 studies).

Regarding studies using repeated tDCS sessions, we found: a series of 30 musicians with right-hand primary focal dystonia who received 10 sessions of tDCS delivered over the both M1 regions (C3/C4, the cathode being on the left side), combined with a rehabilitative intervention (sensory motor retuning therapy) for 2 weeks and who experienced a more significant improvement of their dystonia severity score after active than sham tDCS (Rosset-Llobet et al., 2015). Interestingly, another research group reported that single-session protocols of anodal or cathodal tDCS of left M1 (with right supraorbital cathode) failed to produce any beneficial effects in patients with musician's dystonia (Buttkus et al., 2010, 2011), whereas simultaneous bihemispheric
("dual") stimulation of M1 (left cathode + right anode) improved the rhythmic accuracy of sequential finger movements in such patients, up to 4 days after intervention (Furuya et al., 2014);

- a series of 8 patients with writer's cramp who underwent 3 tDCS sessions in one week, with the cathode placed over the TMS motor hotspot and the anode on the contralateral mastoid (Benninger et al., 2011). The stimulation did not produce beneficial effects on disability and severity of dystonic symptoms and failed to restore normal kinematics of handwriting and cortical inhibition. Sham-stimulated patients even reported a significantly better improvement contrasting with the subjective worsening with tDCS in some patients, which remained unexplained and constituted the reason for the premature termination of the study;

- the case of one patient with cervical dystonia who was not improved by five consecutive daily sessions of tDCS with the cathode at C4 and the anode at P3, according to the international 10/20 EEG electrode system (Angelakis et al., 2013);

- another case of one patient with cervical dystonia who benefitted on dystonia symptoms and quality of life from a sequential protocol of anodal tDCS of the both cerebellar hemispheres and the right M1 performed during 20 sessions over 12 weeks after a botulinum toxin injection (Bradnam et al., 2014);

- and finally a series of 9 young people with mainly generalized dystonia secondary to cerebral palsy who did not respond to a clinically meaningful extent to a 5-day protocol of either cathodal or anodal tDCS of M1 (2 mA, 9 minutes per day) (Bhanpuri et al., 2015).

All these studies are not-replicated studies, based on small sample sizes, excepting the study published by Rosset-Llobet et al. (2015). Therefore, no recommendation can be drawn from these results regarding the efficacy of any tDCS protocol on dystonia symptoms.

6. Motor stroke

The recovery of motor function after stroke is one of the most important issues addressed in neurorehabilitation medicine. The rationale of cortical stimulation following stroke is to promote adaptive neuroplasticity, taking place in the perilesional region as well as in the homologous area of the contralesional hemisphere. Functional magnetic resonance imaging (fMRI) studies have revealed an initial decrease in ipsilesional activity in the first 3 days after stroke followed by an increase in activity in homologous contralesional as well as perilesional areas after 10 days (Grefkes and Ward, 2014). Contralesional activity returns close to normal values when motor function improves but remains elevated when significant clinical impairment persists. The initially reduced ipsilesional activity is associated with a decreased ipsilesional corticospinal excitability as indicated by increased motor threshold and decreased MEP amplitude when tested by TMS. This leads to a contralesional increase in corticospinal excitability due to diminished interhemispheric inhibition from ipsilesional to contralesional M1, which in turn increases interhemispheric inhibition from contralesional to
ipsilesional M1 and further decreases ipsilesional corticospinal excitability (Traversa et al., 1998; Murase et al., 2004). Thus, tDCS therapy aims at either increasing ipsilesional M1 excitability or decreasing contralesional M1 excitability, or both at the same time via bihemispheric tDCS. Most tDCS studies targeted the hand/arm M1 representation, but some studies specifically addressed post-stroke swallowing or lower limb dysfunction. Studies differed in stimulation parameters (tDCS intensity, polarity and duration), in the number of sessions applied, the duration of follow-up, and the tasks performed for outcome evaluation. Finally, tDCS can be used alone or combined in different sequence order with physical or other non-pharmacological rehabilitation therapies.

A PubMed research (keywords: tDCS AND motor AND stroke) identified 261 papers, including 68 original clinical trials and 1,032 patients. The majority of these studies targeted upper limb function. As for the other indications, we excluded from the analysis all studies that do not have a control condition using sham tDCS, a sample size with at least 10 patients receiving active tDCS, and a design consisting of repeated daily sessions. According to these criteria, we retained only 7 studies for making recommendations on the value of tDCS to improve post-stroke motor dysfunction (Table 3).

Two large RCTs, fulfilling the criteria for class I, did not show any significant clinical effect of tDCS on motor function recovery (Hesse et al., 2011; Rossi et al., 2013). Presumably, in one study (Hesse et al., 2011), this was due to the inclusion of patients with cortical stroke and severe motor weakness, who are less susceptible to improve after cortical stimulation (Ameli et al., 2009). The second negative study (Rossi et al., 2013) concerned the immediate acute phase (two days after stroke) by applying anodal tDCS to the ipsilesional M1, whose dramatic reduction in activity in this period (Grefkes and Ward, 2014) may likely not be overcome by tDCS.

The other controlled studies retained in our analysis were of class II (Table 3), reporting either positive or negative results in terms of efficacy of tDCS on motor recovery or quality of life. There is heterogeneity in clinical profile of the population, tDCS methodology, or outcome measures, precluding robust conclusions. For example, regarding anodal tDCS of ipsilesional M1 in chronic stroke patients, leg area was targeted and walking measures were performed in one negative study (Geroin et al., 2011), while hand area was targeted in two positive studies, one showing improvement in quality of life but not in motor performance (Viana et al., 2014) and the other showing improvement in some motor tests but not all (Allman et al., 2016). Overall, the level of evidence remains insufficient to make any recommendation regarding either anodal tDCS of ipsilesional M1 or cathodal tDCS of contralesional M1 in populations of patients at various time after stroke.

A previous meta-analysis of anodal tDCS of ipsilesional M1 in chronic stroke patients, including trials with smaller sample sizes and based on single tDCS sessions, showed small to moderate effect sizes for the improvement of upper limb function when compared with baseline or sham tDCS, respectively (Butler et al., 2013). The same year, another meta-analysis (Elsner et al., 2013b) including all types of tDCS protocols, found evidence of an effect in favour of tDCS on
activities of daily living (primary outcome) at follow-up but not at the end of the intervention phase in stroke patients. The reverse was observed for upper limb motor function, but evidence was of low quality when only studies with low risk of bias were considered. This meta-analysis was updated three years later (Elsner et al., 2016c): the moderate beneficial effect on activities of daily living was found both at the end of the intervention and at follow-up, whereas no more evidence for upper limb motor function improvement could be observed. Another meta-analysis including multiple- and single-session studies combining tDCS and motor training of the upper extremities showed moderate long-term effects on motor learning after either anodal tDCS of ipsilesional M1, cathodal tDCS of contralesional M1, or bihemispheric stimulation of M1 in the postacute or chronic stage of poststroke recovery (Kang et al., 2016). A fourth meta-analysis, including only RCTs with a multiple-session design, showed only small non-significant beneficial effects of therapeutic strategies combining tDCS and rehabilitation programs for poststroke upper extremity recovery (Tedesco Triccas et al., 2016). A fifth meta-analysis determined whether the outcome of tDCS of M1 in poststroke motor recovery has a dose-response relationship (Chhatbar et al., 2016). This study included 8 sham-controlled RCTs based on repeated tDCS sessions (≥5) including assessment on the Fugl-Meyer Assessment (FMA) scale, concerning a total of 213 stroke patients. This study showed a moderate effect of active stimulation on FMA scores with greater effects of bihemispheric stimulation and in chronic stroke patients. It further revealed a positive dose-response relationship for current and charge density as well as a negative dose-response relationship for electrode size.

Additionally, some results obtained with single tDCS sessions are worth being discussed, although they should be considered as "proof-of-concept" studies, based on immediate or short-lasting effects rather than "therapeutical trials" aimed at producing sustained effects, which could be clinically relevant. In chronic stroke, several single-session studies showed beneficial effects of contralesional cathodal stimulation of M1 on motor performance of the paretic hand, e.g., assessed by the Jebsen-Taylor hand function test (JTT) (10 patients investigated 1-16 months after stroke in Mahmoudi et al., 2011) or a finger movement task (12 patients investigated 12-64 months after stroke in Zimerman et al., 2012). Short-lasting motor performance improvement was also reported after a single session of ipsilesional anodal stimulation of M1, e.g., regarding pinch force and motor response time of the paretic hand (11 patients investigated 18-107 months after stroke in Hummel et al., 2006; 13 patients investigated 18-70 months after stroke in Stagg et al., 2012). One study showed that tDCS effects were greater after ipsilesional anodal stimulation of M1 than contralesional cathodal stimulation (Stagg et al., 2012).

A single anodal tDCS session targeted over the leg area of ipsilesional M1 was also found to improve gait and lower limb strength in a series of 11 patients in the postacute phase (1-3 months after stroke) (Sohn et al., 2013). The same montage was used in a multiple-session study, showing some improvement of motor function in the lower limbs, but not of gait performance (Chang et al., 2015). Another study, still using the same tDCS protocol but combined with robot-assisted gait training for
10 sessions in patients with chronic stroke (14-34 months after stroke) did not confirm the additional value of anodal tDCS targeted over the leg area of ipsilesional M1 to improve walking abilities (Geroin et al., 2011). The absence of additional value of tDCS when combined with robot therapy was confirmed for upper extremity rehabilitation, evaluated on FMA scale in a recent study (Triccas et al., 2015). In this study, 18 sessions of ipsilesional anodal tDCS were performed over 8 weeks in a series of 22 patients either in the postacute (2-3 months after stroke) or chronic phase (9-90 months after stroke). Both active and sham tDCS groups included only 6 postacute stroke patients and 5 chronic stroke patients. Because of this heterogeneity and small sample size, the study was not entered into the analysis for grading. These negative results probably revealed a ceiling effect of very thorough rehabilitation programs. Conversely, tDCS was found to be more beneficial than basic functional training for improving upper and lower limb functions in a parallel-arm study of 20 chronic stroke patients receiving ipsilesional anodal tDCS, 5 days a week for 4 weeks (Cha et al., 2014).

All above cited results were obtained with the "return" ("reference") tDCS electrode placed over the contralateral supraorbital region. Conversely, some studies also considered bihemispheric stimulation of M1, with the cathode over the contralesional M1 region and the anode over the lesioned M1. One single-session study showed that bilateral montage was as efficacious as unilateral cathodal or anodal stimulation (Mahmoudi et al., 2011). Also in proof-of-concept studies based on a single session, one group showed the value of bihemispheric M1 tDCS protocol combined with a visuomotor skill learning task (Lefebvre et al., 2012) or precision grip and dexterity tests (Lefebvre et al., 2014) to enhance motor rehabilitation in chronic stroke patients. However, another study did not confirm the value of a single bihemispheric tDCS session on walking abilities in a small series of chronic stroke patients (van Asseldonk and Boonstra, 2016). To our knowledge, there is only one study reporting the clinical impact of a 5-day protocol of bihemispheric M1 tDCS, combined with 60-min occupational therapy, in a series of 20 patients with chronic stroke (5-81 months after stroke) (Lindenberg et al., 2010). In this parallel-arm study, an improvement of motor performance on FMA and Wolf motor function test (WMFT), lasting for one week beyond the time of stimulation, was observed in the 10 patients having received active tDCS. However, in the absence of replicated bihemispheric tDCS studies including repeated sessions, no recommendation can be proposed for this approach. In addition, the superiority of unilateral anodal and cathodal M1 stimulation over bilateral M1 stimulation was only shown regarding the effect of tDCS on motor corticospinal excitability in healthy volunteers or speeding reaction time in chronic stroke patients (O'Shea et al., 2014). The potential impact of bihemispheric M1 tDCS on motor rehabilitation may depend on the microstructural and functional status of transcallosal motor tracts (Lindenberg et al., 2016). However, the concept of a bihemispheric stimulation leading to enhance the excitability of the lesioned motor cortex by the anode and to reduce the excitability of the contralesional hemisphere by the cathode remains to be demonstrated. In particular, a “titration” study is needed for bihemispheric versus monohemispheric...
stimulation of M1 (Batsikadze et al., 2013), to determine the respective impact on motor cortex excitability of these two montages, according to the resulting changes in MEP size, for example.

In the present work, we have differentiated the results according to the time window following stroke, which is known to be a crucial parameter in the focal changes of brain excitability and plasticity. The time of the intervention after stroke onset has very likely significant impact on the efficacy of a given tDCS paradigm. In the acute phase of stroke, a beneficial effect of cathodal tDCS to reduce the inhibitory action exerted by the contralesional M1 onto the lesioned hemisphere should be expected. However, the adaptive role of the contralesional hemisphere in stroke recovery remains controversial (Di Pino et al., 2014). As mentioned above, the classical concept of interhemispheric competitive rivalry supports the objective of decreasing the excitability of the contralesional motor cortex and its ability to further reduce the neural activities in the lesioned hemisphere (Marshall et al., 2000; Ward et al., 2003; Murase et al., 2004). Conversely, some fMRI studies showed a positive correlation between the gain after rehabilitation therapy and the level of cortical activation in the contralesional hemisphere (Cramer et al., 1997; Johansen-Berg et al., 2002a). Furthermore, one study showed that the improvement in motor function and dexterity of stroke patients receiving anodal tDCS over the lesioned motor cortex were associated with an increase in the excitability of the contralesional rather than the ipsilesional hemisphere (Cunningham et al., 2015). In fact, the beneficial contribution of contralesional hemispheric activation in motor performance of the paretic limb might primarily result from the involvement of premotor rather than primary motor areas (Caramia et al., 2000; Johansen-Berg et al., 2002b). In addition, the influence of inhibitory projections from the contralesional brain areas to the spinal cord depends on the considered motor region (either proximal or distal limb muscles), the degree of functional impairment, and the extent of the neuronal damage.

Indeed, the most important factor for predicting brain stimulation efficacy in motor stroke seems to be the integrity of the stimulated cortical region and the corresponding corticospinal tract, as assessed by MEP recordings (Talelli et al., 2006, van Kuijk et al., 2009) or neuroimaging methods (Riley et al., 2011; Bradnam et al., 2012). When the pyramidal motor structures are rather preserved in mildly to moderately affected patients, inhibition of contralesional M1 by cathodal tDCS may promote stroke recovery at least in the post-acute and chronic phases. When neuronal destruction is more extensive in moderately to severely affected patients, descending projections from the contralesional hemisphere contribute to recovery and cathodal tDCS of contralesional M1 may lead to further clinical deterioration (Bradnam et al., 2012). In this case, reducing perilesional activity could be a better strategy (Wu et al., 2013). Another factor involved in the outcome of tDCS therapy appears to be the level of residual GABA activity in the ipsilesional hemisphere. Actually, a greater motor improvement results from anodal tDCS of ipsilesional M1 in patients exhibiting higher GABA activity in the ipsilesional cortex (O'Shea et al., 2014). A bimodal balance-recovery model, linking interhemispheric competition and functional recovery to the structural “neuronal reserve” spared by the lesion, was
recently proposed to enable NIBS protocols to be tailored to the needs of individual patients (Di Pino et al., 2014).

To conclude, the possibility of promoting motor stroke recovery by tDCS still needs to be demonstrated in large, multicentric RCTs consisting of repeated sessions with prolonged follow-up. According to the currently published data, we can identify only some trends in predicting tDCS efficacy. The combination of tDCS with other therapies (e.g., virtual reality training (VRTX) (Lee and Chun, 2014; Viana et al., 2014), occupational therapy (Nair et al., 2011; Zheng and Schlaug, 2015), robot-assisted training (Ochi et al., 2013; Picelli et al., 2015), or constraint-induced movement therapy (Bolognini et al., 2011; Cunningham et al., 2015; Rocha et al., 2016)) could result in synergistic effects to bring the effect of tDCS at a clinically meaningful level. This point is of particular importance and one of the main challenges in daily practice will be to optimize the combined approach of tDCS and rehabilitation therapies to produce maximum synergy in both directions, i.e. increasing the efficacy of tDCS through the action of rehabilitation therapies and vice versa.

In the future, experimental designs should also take into account the influence of stimulation parameters (site, intensity, duration), the phase of stroke (acute, post-acute, chronic), the degree of clinical impairment, and more importantly, the location and extent of stroke lesion. A proper selection of patients is needed, substantiated by a careful assessment of various anatomo-functional parameters at baseline, to avoid a negative impact of unsuccessful trials on the development of cortical stimulation strategies for treating stroke. As examples of failure, there are trials that included patients with severe cortical stroke unable to respond to such treatment (e.g., Hesse et al., 2011 for tDCS) or protocols that did not take into account stroke-induced plasticity to design stimulation location and settings (e.g., Harvey et al., 2009 for the EVEREST trial of implanted epidural cortical stimulation). Thus, the plastic changes in motor cortical representation and the integrity of the corticospinal tract should be carefully investigated by means of motor cortical mapping using navigated TMS (Ahmad et al., 2014; Mellerio et al., 2014) and functional neuroimaging techniques, such as diffusion tensor imaging tractography or analyses of resting-state or task-dependent motor network connectivity.

There are potentially other interesting indications of tDCS in the context of stroke, such as swallowing dysfunction (Kumar et al., 2011; Yang et al., 2012; Shigematsu et al., 2013) or spasticity (Hesse et al., 2011; Ochi et al., 2013; Vandermeeren et al., 2013; Wu et al., 2013; Lee and Chun, 2014; Viana et al., 2014). The issue of dysphagia was not further addressed in this work, since only one study including more than 10 patients receiving the active treatment has been reported so far (Shigematsu et al., 2013). Regarding spasticity, tDCS was mostly combined with other therapeutic strategies (VRTX, physical therapy, robot-assisted training) as mentioned above. There were three large-scale sham-controlled studies: two are presented in Table 3 (Hesse et al., 2011; Viana et al., 2014) and one was a non-replicated study of cathodal tDCS of the contralesional motor cortex in patients in the chronic stroke phase (Wu et al., 2013). Reviews and meta-analyses can be found
elsewhere on tDCS and dysphagia (Doeltgen et al., 2015; Yang et al., 2015; Pisegna et al., 2016) or spasticity (Elsner et al., 2016a).

Beyond motor impairment, stroke may also affect other neurological functions, such as speech and language production or awareness of the contralesional body or environment. A deficit in the first of these functions refers to aphasia, which will be discussed in the next chapter, whereas a deficit in the second of these functions refers to hemispatial neglect. A PubMed research (keywords: tDCS AND neglect AND stroke) identified 17 papers, including 6 original clinical trials and 41 patients. These studies included only one to ten patients, with a certain methodological heterogeneity, therefore precluding any recommendation in this clinical condition.

Please insert TABLE 3

7. Aphasia

Aphasia is a highly disabling language disorder frequently caused by a left-lateralized hemispheric stroke (Laska et al., 2001). Even if traditional linguistic-based therapies have been proved to induce an adequate clinical improvement (Basso et al., 2011; Code and Petheram, 2011; Brady et al., 2012), a large percentage of patients are left with some degree of language impairment (Lazar et al., 2010). It has been suggested that one of the key predictors for positive language outcomes is treatment intensity (i.e., more than 5 h of speech therapy per week) (Bhogal et al., 2003). Therefore, effective treatment approaches that might be used as an adjunct to common speech and language therapies are urgently needed to maximize the recovery process in aphasic patients. The recent application of NIBS techniques to language rehabilitation, such as tDCS, has already provided positive results (Monti et al., 2013). Generally, interhemispheric competition between the residual language areas in the damaged left hemisphere and the intact right hemisphere underlies the basic concept for tDCS language treatment (Kiran, 2012). In particular, it has been proposed that in patients with left hemispheric damage, the homotopic contralateral right hemispheric areas may be in a state of abnormally high activation and may exert an inhibitory effect over the stroke-affected hemisphere (Belin et al., 1996; Murase et al., 2004). Thus, a language improvement may be possible either by increasing the output of the perilesional left hemisphere through excitatory (anodal) tDCS, by decreasing the inhibition from the intact right hemisphere by applying inhibitory (cathodal) tDCS over the contralesional cortex, or both.

A PubMed search (keywords: tDCS AND aphasia) identified 102 papers, including 35 original clinical studies and 431 patients. Studies mainly concerned patients with nonfluential poststroke aphasia, while there are three studies on primary progressive aphasia (Wang et al., 2013; Cotelli et al., 2014b; Tsapkini et al., 2014), which will be not analyzed further, because of the low volume of reported data and the methodological heterogeneity of those studies. Regarding nonfluential poststroke aphasia, most
studies were single case reports or small series of patients, including less than 10 patients receiving active stimulation, or based on single tDCS sessions (Class III or IV studies). Therefore, only 5 studies were retained for analysis regarding tDCS effects on nonfluent aphasics (Table 4). Anodal tDCS was applied over the Broca’s area (left inferior frontal gyrus) in four studies and over the Wernicke’s area (left postero-superior temporal gyrus) in two studies. However, we have to mention that the first tDCS work on aphasia showed that only cathodal tDCS (and not anodal tDCS) over a left frontotemporal region significantly improved the accuracy of the picture naming task in 8 chronic non-fluent post-stroke aphasic patients (Monti et al., 2008).

Regarding anodal tDCS applied over Broca’s area, several sham-controlled studies issued from the same group (Fiori et al., 2013; Marangolo et al., 2013a,c, 2014a; Campana et al., 2015) and performed in series of 7 to 20 patients with nonfluent poststroke aphasia were in favour of a significant improvement of speech and language performance at the end of a protocol of 5-10 sessions of active tDCS. Conversely, two sham-controlled studies from another group (Polanowska et al., 2013a,b) did not find significant difference between active and sham conditions in language recovery after 15 sessions of tDCS performed in 24 nonfluent aphasics. These two sets of studies present several differences, including the post-stroke phase in which patients were included: either the postacute phase (2 months in average) (Polanowska et al., 2013a,b) or the chronic phase (ranging from 6 months to 7 years) (Fiori et al., 2013; Marangolo et al., 2013a,c; Campana et al., 2015). Anyway, no recommendation can be drawn from these results.

There are several other studies based on single sessions or smaller sample size with the same tDCS montage. In the study published by Volpato et al. (2013), 2 nonfluent and 6 fluent aphasics at chronic stage (6-126 months after stroke) did not benefit from 10 sessions of anodal tDCS over Broca’s area in object and action naming accuracy. In this latter study, tDCS was applied as a "single" therapy. In contrast, various studies showed the potential value of coupling multiple sessions of anodal tDCS over Broca’s area with concurrent language or conversational therapy (Baker et al., 2010; Marangolo et al., 2011, 2013a,c, 2014a; Fiori et al., 2013; Campana et al., 2015). Except the study by Baker et al. (2010), all other studies were issued from the same group, showing significant tDCS-induced changes not only in naming performance but also in the recovery from articulatory disturbances and speech production deficits. In the study of Baker et al. (2010), 4 nonfluent and 6 fluent aphasics at a chronic stage (10-242 months after stroke) received 5 sessions of anodal tDCS over the left frontal cortex while performing a computerized anomia treatment. In this study, fMRI investigation was performed during an overt naming task in each individual to position the active anode over structurally intact frontal cortex. The tDCS protocol improved naming accuracy at least during one-week follow-up. One of the two non-fluent aphasics who received stimulation specifically over the Broca’s area benefited from the treatment.

Regarding anodal tDCS applied over Wernicke’s area, three studies found beneficial results of a protocol of 5 sessions of tDCS coupled with naming treatment in 12 patients with various types of
aphasia in the postacute phase (3-6 months after stroke) (Wu et al., 2015) and in smaller groups of 3 nonfluent (Fiori et al., 2011) and 8 fluent (Fridriksson et al., 2011) aphasics at the chronic phase, with the improvement lasting up to 3 weeks after the end of the intervention. In contrast, this montage did not provide any significant improvement in auditory comprehension and naming in a series of 7 patients with various types of aphasia in the postacute phase (about one month after stroke) (You et al., 2011). Again, these results preclude any recommendation for anodal tDCS over the left hemisphere.

However, according to the concept of interhemispheric rivalry as the origin of poststroke functional changes, other tDCS studies aimed at stimulating cathodally the homologous regions of Broca’s and Wernicke’s areas in the right hemisphere.

Cathodal tDCS of the right homologous of Broca’s area was performed in four studies (Jung et al., 2011; Kang et al., 2011; Vines et al., 2011; Cipollari et al., 2015), based on multiple session (3-10) protocols in patients at either postacute or chronic stage. Two of these studies showed improvement in verbal fluency but included only 6 nonfluent aphasics (Vines et al., 2011; Cipollari et al., 2015), while the other two studies were performed on respectively 10 (Kang et al., 2011) and 37 patients (Jung et al., 2011), the latter study being based on an open-label protocol. In the study by Kang et al. (2011), each patient received a daily session of cathodal tDCS for 5 consecutive days in a randomized crossover manner coupled with conventional word-retrieval training. Significantly improved picture naming was observed at 1 hour following the last active tDCS session but no changes were observed after sham tDCS. Some improvement was also reported after cathodal tDCS of the right homologous of Wernicke’s area in a single case (Cheney et al., 2013) and in a series of 7 patients with various types of aphasia in the postacute phase (You et al., 2011). In contrast, this montage did not provide any significant improvement in 12 chronic aphasics (Floel et al., 2012). Therefore, these results do not allow any recommendation to be proposed.

Another approach was to combine both anodal tDCS of the left inferior frontal gyrus and cathodal tDCS of the right inferior frontal gyrus in a bihemispheric (“dual”) tDCS strategy. The first proof-of-concept was provided by an open-labelled single-session study performed in 6 nonfluent and 5 fluent aphasics at a chronic stage (Lee et al., 2013). Marangolo et al. (2013b, 2014b, 2016) and Cipollari et al. (2015) extended this result in a series of 6-9 nonfluent aphasics at a chronic stage receiving 10-15 sessions of bihemispheric tDCS combined with speech and language therapy. Improvement in picture description, noun and verb naming, word reading and repetition accuracy, and response time lasted up to one week after the intervention. However, these results still remain to be reproduced on a larger sample size by another independent team. It is interesting to note that a similar strategy of bihemispheric (“dual”) stimulation was successfully applied using high-frequency rTMS over Broca’s area combined with low-frequency rTMS over its right homologous to promote poststroke rehabilitation of aphasia (Khedr et al., 2014a).
We also have to mention that a bihemispheric tDCS strategy targeted on the DLPFC (and not the inferior frontal gyrus) was successfully applied in two open-labelled case series of 1-4 patients with chronic nonfluent aphasia (Manenti et al., 2015; Valiengo et al., 2016). Finally, one group chose the original strategy of stimulating the motor cortex combined with computer-assisted language therapy to improve recovery outcome in poststroke aphasia (Meinzer et al., 2016). In a recently reported trial including 26 patients with various types of poststroke aphasia, anodal tDCS administered for 2 weeks over the left M1 (C3 site, with the cathode over the right supraorbital region) was found to produce a beneficial effect on naming ability. Improvement after active vs. sham procedure was especially larger for trained items during follow-up (6 months after stimulation) and regarding transfer to untrained items (Meinzer et al., 2016). However, the value of this approach remains to be replicated by another research group.

To conclude, some beneficial results were reported using anodal tDCS over Broca's and Wernicke's areas, cathodal tDCS over the right homologue of Broca's area, or a bihemispheric stimulation of both inferior frontal gyri. However, the level of evidence is not sufficient to ensure the efficacy or the therapeutic potential of any of these protocols in the rehabilitation of poststroke aphasics. A similar conclusion was reported in a meta-analysis issued in 2013 and updated in 2015 (Elsner et al., 2013a, 2015). Moreover, direct comparisons between these different montages and approaches are lacking. Therefore, further studies are needed to identify which parameters of stimulation (e.g., anodal or cathodal monohemispheric or bihemispheric stimulation, stimulation intensity, session duration, interval between sessions...) can maximize tDCS effect on the recovery process. Most studies are case reports or have small sample size and some of them did not monitor the benefits obtained over time during and beyond the period of stimulation. Large, multicentric RCTs are needed to replicate the results and to assess different outcome measures, such as standardized tests of everyday communication abilities. If promoting effects of tDCS on stroke recovery are confirmed in larger samples of aphasic patients, the parameters of stimulation to use in daily practice should be clearly determined, as well as the optimal time after stroke to initiate the treatment, the type of aphasia to manage (nonfluent, fluent, or both) and the concomitant speech and language therapy to perform.

Please insert TABLE 4

8. Multiple sclerosis

Multiple sclerosis (MS) is one of the most common neurological diseases and a serious cause of disability in young adults. Its natural course is characterized by recurrent relapses or progressive functional decline. With disease evolution, patients could accumulate several neurological dysfunctions or disease-related complications including motor deficit, fatigue, tremor, spasticity, sensory disturbances, pain, genital or urinary symptoms, and psychiatric or cognitive disorders. Many
of these clinical aspects are difficult to manage, being resistant to pharmacological treatments. Thus, dealing with MS complications constitutes a real challenge to every person in charge, making new approaches much needed in this domain. The effects of tDCS have been assessed on various symptoms in MS patients. A PubMed search (keywords: tDCS AND multiple sclerosis) identified 19 papers, including 11 original clinical studies and 205 patients. The topic of these 11 studies was as follows:

- fatigue (Ferrucci et al., 2014; Saiote et al., 2014; Tecchio et al., 2014, 2015; Ayache et al., 2016);
- cognition, attention and executive function (Ayache et al., 2016; Mattioli et al., 2016);
- motor performance (Cuypers et al., 2013; Meesen et al., 2014);
- spasticity (Iodice et al., 2015);
- pain or tactile sensory deficit (Mori et al., 2010, 2013; Ayache et al., 2016).

To date, the main MS domain investigated in tDCS trials concerned fatigue, a very frequent symptom in MS patients, with a complex pathophysiology, mixing pathological changes related to the disease itself and various comorbidities which may be accessible to NIBS techniques (Palm et al., 2014a; Chalah et al., 2015). Protocols varied greatly between studies, including the cortical target, which was either the bilateral motor cortex (Ferrucci et al., 2014), the bilateral sensory or sensory-motor cortex (Tecchio et al., 2014, 2015), or the left DLPFC (Saiote et al., 2014; Ayache et al., 2016).

Data provided by motor or sensory cortex tDCS are summarized in Table 5, showing a mean reduction of 28% of the score on the modified fatigue impact scale (mFIS) after active tDCS, whatever the montage and the study. However, given the methodological heterogeneity of the two types of study (Ferrucci et al., 2014 vs. Tecchio et al., 2014, 2015), no recommendation can be made. Regarding anodal tDCS of the left DLPFC (F3), no significant effect was observed after either 3 daily sessions performed in 8 patients with relapsing-remitting (RR) or progressive MS (Ayache et al., 2016) or 5 daily sessions performed in 13 patients with RR-MS (Saiote et al., 2014). However, in the latter study, the responders to tDCS were characterized by a higher prefrontal lesion load in brain MRI analysis, which might favour tDCS targeting over the lesioned areas in MS.

Regarding cognition and executive function, Mattioli et al. (2016) reported in 20 MS patients (10 in the active arm and 10 in the sham arm) that 10 daily sessions of anodal tDCS of the left DLPFC could improve the impact of cognitive training (performed concurrently with tDCS) on attention and speed of information processing. The effects were found to persist up to 6 months after the intervention. These effects on attention were not replicated in another study of 16 MS patients (8 in the active arm and 8 in the sham arm) (Ayache et al., 2016).

Regarding motor performance, Cuypers et al. (2013) showed that a single session of anodal tDCS of M1, contralateral to the most impaired hand, increased motor corticospinal output and strength in MS patients. Therefore, the authors suggested that tDCS could be considered in motor training protocols for the rehabilitation of MS patients. However, Meesen et al. (2014) tested the add-
on value of a single session of anodal tDCS of M1, contralateral to the most impaired hand, on motor training in MS patients. They did not find any difference between active and sham conditions, especially regarding finger tapping test. These single-session results should be interpreted with caution, since multiple sessions are probably required to bring tDCS effects to a clinically meaningful level.

Regarding spasticity, Iodice et al. (2015) showed in a sham-controlled parallel-arm study of 20 RR-MS patients that a 5-day protocol of anodal tDCS delivered to M1 of the most affected side produced no significant impact on the modified Ashworth scale (MAS) or other scales relative to MS spasticity or walking abilities.

Finally, three studies with a parallel-arm design investigated the effect of tDCS on sensory aspects of MS. The first study consisted of 5 sessions of anodal tDCS applied over the motor cortex (C3/C4 in the 10-20 system of EEG electrode placement) contralateral to the painful somatic area in 19 RR-MS patients (10 in the active arm and 9 in the sham arm) (Mori et al., 2010). A significant pain relief was observed after active tDCS compared to sham tDCS, while there was no change in depression or anxiety scores. In the second study, the same authors performed 5 sessions of anodal tDCS 2 cm posteriorly to the C3/C4 position in order to stimulate S1, contralateral to the hypoesthetic upper limb of 20 RR-MS patients (10 in the active arm and 10 in the sham arm) (Mori et al., 2013). They found a significant improvement of tactile discriminatory thresholds and sensation scores after anodal tDCS compared to sham stimulation, again without any impact on depression scores (Mori et al., 2013). Finally, in a third study including 16 MS patients (8 in the active arm and 8 in the sham arm), three consecutive daily sessions of anodal tDCS applied to the left DLPFC were found to produce significant analgesic effects compared to a sham condition, without concomitant improvement on mood, fatigue, or attention (Ayache et al., 2016).

In summary, tDCS studies in the MS domain are sparse and results are still preliminary and heterogeneous regarding the methods used and the symptoms to treat. Therefore, no recommendations can be made.

Please insert TABLE 5

9. Epilepsy

Although NIBS techniques aimed at modifying cortical excitability, the study of their therapeutic potential in epilepsy remains underdeveloped. This is the case for rTMS (Lefaucheur et al., 2014) and even more for tDCS. Actually, a PubMed search (keywords: tDCS AND Epilepsy) identified 65 papers, including only 10 original clinical studies and 147 patients. Moreover, 5 of these studies were case reports of 1 to 5 patients, while 5 sham-controlled studies with crossover or parallel-arm design included 12 to 37 patients (Fregni et al., 2006g; Auvichayapat et al., 2013; Liu et al., 2015; San-Juan
et al., 2016; Tekturk et al., 2016b). Two of these studies were based on a single tDCS session (Fregni et al., 2006g; Auvichayapat et al., 2013), while another study (Liu et al., 2015) explored the safety and antidepressant efficacy of a multiple (5-day) tDCS session protocol in a series of epileptic patients, but did not address the question of the antiepileptic potential of tDCS.

In Fregni et al. (2006g), tDCS was targeted over the epileptogenic focus in 19 patients with focal epilepsy due to cortical dysplasia. A single session of active cathodal tDCS delivered at 1 mA for 20 min and performed in 10 of these patients, led to a reduction in epileptic discharge frequency, with a trend towards a decrease in the number of clinical seizures. The remaining 9 patients did not benefit from sham tDCS. In Auvichayapat et al. (2013), cathodal tDCS was also applied at 1 mA as a single session of 20 min over the seizure focus in a series of 29 children with focal epilepsy (29 patients in the active tDCS group and 7 patients in the sham tDCS group). A reduction in epileptic discharge frequency was observed up to 48 hours after active tDCS, but the final result was clinically negligible.

In Liu et al. (2015), anodal tDCS (2 mA) was applied for 20 min over the left DLPFC (F3), with the cathode over the right supraorbital area, for 5 consecutive days, according to a depression therapy protocol. Thirty-three patients with temporal lobe epilepsy completed the study (21 patients in the active tDCS group and 12 patients in the sham tDCS group). Beyond the antidepressant effects, active tDCS did not increase seizure or interictal discharge frequency. This study provides evidence for the safety of anodal tDCS of left DLPFC in epileptic patients, but was not intended to investigate antiepileptic therapy.

Few case reports or small sample studies have shown the value of repeated cathodal tDCS sessions applied over the epileptic focus to reduce epileptic EEG activity or the number of seizures, e.g. 4-5 tDCS sessions in 2-5 patients with Rasmussen's encephalitis (San-Juan et al., 2011; Tekturk et al., 2016a) and 2 tDCS sessions in 2 patients with focal epilepsy (Assenza et al., 2014). The effect of repeated cathodal tDCS sessions on seizure frequency was recently reported in larger populations of patients with mesial temporal lobe epilepsy and hippocampal sclerosis (San-Juan et al., 2016; Tekturk et al., 2016b). In the first study of 28 patients with a parallel-arm design (20 patients in the active arm and only 8 patients in the sham arm) (San-Juan et al., 2016), the cathode was positioned over the most active area of interictal epileptiform discharges defined on scalp EEG. In the second study of 12 patients with a crossover design (Tekturk et al., 2016b), the cathode was placed over the temporal region (T3 or T4), contralateral to the most affected side. In both studies, the anode was placed over the contralateral supraorbital region and tDCS was delivered at 2mA for 30 minutes in 3 to 5 sessions performed on consecutive days. The mean seizure frequency significantly decreased after active but not sham tDCS in both studies, even associated with a reduction of interictal epileptiform discharges on scalp EEG immediately after tDCS (San-Juan et al., 2016). However, the comparability of the procedure used in these two studies remains to be determined, especially regarding cathode location and therefore, no recommendation can be made about the potential efficacy of tDCS in the treatment of any type of epilepsy.
10. Disorders of consciousness

Disorders of consciousness, such as minimally conscious state (MCS) and vegetative state (VS), are highly challenging clinical conditions for treatment. A PubMed search (keywords: tDCS AND vegetative state OR disorders of consciousness) identified 23 papers, including 4 original clinical studies and 79 patients. These four studies included 10, 30, 25, and 14 patients, respectively (Angelakis et al., 2014; Thibaut et al., 2014; Naroe et al., 2015, 2016). The left DLPFC was targeted with anodal tDCS applied for 20 min at 2 mA and for 5 days in a first study based on a limited number of patients (3 patients in MCS and 7 patients in VS) (Angelakis et al., 2014) and only as a single session for the second study, which included a larger sample size (30 patients in MCS and 25 patients in VS) (Thibaut et al., 2014). Both studies, from independent groups, arrived at a similar conclusion: patients with MCS but not those with persistent VS or coma can benefit from anodal tDCS of the left DLPFC, combined with cathodal tDCS of the right homologue area (Thibaut et al., 2014) or the right orbitofrontal cortex (Angelakis et al., 2014).

Another group delivered single sessions of anodal tDCS over the orbitofrontal cortex, the DLPFC, or a parieto-occipital region in 25 and 14 patients affected by various disorders of consciousness (Naro et al., 2015, 2016). More particularly, these authors intended to show that the response to tDCS could demonstrate the persistence of residual functional connectivity within large-scale brain networks in patients with MCS or more severe disorders of consciousness.

In conclusion, some beneficial results of tDCS protocols have been shown in patients with disorders of consciousness, especially targeting the left DLPFC in MCS. However, the reported data are very preliminary, obtained in small samples, with heterogeneous outcome measures, including either clinical or functional connectivity variables. Therefore, no recommendation can be made, especially regarding the number of sessions and the amount and clinical profile of the patients to treat.

11. Alzheimer’s disease

Since its updating in the early 2000s, tDCS has been widely used in neuropsychological studies to act on cognitive and behavioural features, e.g., attention, memory and working memory, computation, decision-making, and so on, in either healthy humans or patients with various neuropsychiatric diseases (Shin et al., 2015; Hill et al., 2016). Therefore, there are a lot of expectations regarding the therapeutic potential of tDCS to modulate psychological processes in cognitive disorders, such as Alzheimer’s disease (AD). A PubMed search (keywords: tDCS AND Alzheimer’s disease) identified 46 papers, including 9 original clinical studies and 189 patients. Among these 9 studies, there are 1 case report, 1 study on patients with mild cognitive impairment (MCI), and 7 studies that included 10
to 40 AD patients (Ferrucci et al., 2008a; Boggio et al., 2009a, 2012; Cotelli et al., 2014a; Khedr et al., 2014b; Suemoto et al., 2014; Bystad et al., 2016).

The main target explored was the left DLPFC, which was intended to be activated by anodal tDCS to improve cognitive performance. In this domain, the four published studies showed different primary outcome assessments that were a visual recognition memory task (Boggio et al., 2009a), a face-name association task (Cotelli et al., 2014a), an apathy score (Suemoto et al., 2014), or the Mini Mental State Examination (MMSE) (Khedr et al., 2014b). The initial single-session study showed memory improvement in 10 patients having received active anodal tDCS of the left DLPFC (Boggio et al., 2009a). Three more recent sham-controlled studies were retained for analysis (Table 6), including 11 to 20 AD patients having received the active condition in a protocol comprising several tDCS sessions. No safety problems emerged in these studies, in which the left DLPFC was targeted with anodal tDCS applied for 20-25 min at 2 mA for 6-10 sessions performed within two weeks. One study also considered cathodal stimulation of the left DLPFC (Khedr et al., 2014b), while another study explored the value of tDCS combined with memory training (Cotelli et al., 2014a). Although one study showed a significant cognitive improvement on MMSE after both anodal and cathodal tDCS of the left DLPFC, lasting for up to 2 months after the intervention (Khedr et al., 2014b), the other two studies were negative. One study showed that active tDCS had no additional value compared to sham tDCS on the improvement in cognitive performance provided by memory training (Cotelli et al., 2014a). The other study, the largest sham-controlled RCT published to date in this domain (Suemoto et al., 2014), which was adequately powered for disclosing minimally clinically significant differences after the intervention, did not show any significant effect of the active anodal tDCS condition targeting the left DLPFC. Therefore, no recommendation can be currently made regarding the efficacy of anodal tDCS of the left DLPFC to improve cognitive performance in AD patients.

Besides the left DLPFC, the other studies aimed at targeting a temporo-parietal region bilaterally with anodal tDCS, placing one anode over each hemisphere and a cathode at an extracephalic location (deltoid muscle). The stimulation was applied for 15-30 min at 2 mA during repeated daily sessions for 5 days in one study (Boggio et al., 2012) but only in a single session in two other studies that included 10 AD patients each (Ferrucci et al., 2008a; Boggio et al., 2009a). The primary aim was to improve recognition memory. The initial single-session studies showed beneficial effect on word or visual recognition memory, but this result was not confirmed by a subsequent multiple-session RCT (Boggio et al., 2012). In this latter crossover study, including 15 AD patients, visual recognition memory was improved after active vs. sham anodal tDCS of the both temporal cortices at one month after the intervention, but the statistical interaction “time x group” was not significant. Therefore, no recommendation can be currently made regarding the efficacy of bilateral anodal tDCS of the temporo-parietal cortex to improve cognitive performance in AD patients.

Please insert TABLE 6
12. Tinnitus

Tinnitus is a phantom perception of a sound in the absence of a corresponding external sound source and occurs in 5–15% of the population (Axelsson and Ringdahl, 1989; Heller, 2003; Gallus et al., 2015). Whereas many patients can habituate to this sound, quality of life is severely disrupted in about 25% of the patients who cannot cope with the tinnitus (Axelsson and Ringdahl, 1989). In these patients, tinnitus is frequently associated with anxiety, depression, cognitive impairment, and sleep disturbances.

Although numerous treatments have been introduced, evidence for an uniformly successful treatment is lacking (Langguth et al., 2013). The lack of efficient therapies for tinnitus partly originates from the heterogeneity of tinnitus and an incomplete understanding of the pathophysiology of the different forms of tinnitus (Elgoyhen et al., 2015). In most cases, tinnitus is triggered by hearing loss or abnormalities in the inner ear or the peripheral auditory pathway. This results in neuroplastic changes in the central nervous system encompassing a wide network of auditory and non-auditory brain areas (De Ridder et al., 2014). However, some forms of tinnitus appear to be unrelated to hearing loss, which probably reflect a different pathophysiological mechanism and require therefore perhaps a different strategy of neuromodulation (Vanneste and De Ridder 2015).

Over the last decade, tDCS has been used for elucidating the involvement of specific brain networks in tinnitus pathophysiology and has also been investigated as a novel therapeutic approach for tinnitus patients (Langguth et al., 2013). A PubMed search (keywords: tDCS AND tinnitus) identified 48 papers, including 23 original clinical studies and 2,335 patients, but the included patients were largely redundant among studies. In addition, most of these studies have investigated the immediate or short-term effect of single tDCS sessions on tinnitus perception. Longer lasting effects of multiple tDCS sessions were only assessed in 9 studies involving 172 patients treated with active tDCS. Finally, a variety of tDCS montages have been evaluated, targeting either the auditory cortex or the DLPFC or both.

From literature data, only 4 original sham-controlled studies including at least 10 patients who received active tDCS for multiple sessions were retained for analysis (Table 7). All these studies concerned anodal tDCS of the left temporal or temporoparietal (auditory) cortex. Neuroimaging has demonstrated abnormalities of oscillatory brain activity, connectivity and metabolism in the auditory cortex of tinnitus patients (De Ridder et al., 2014). These changes in the auditory cortical area led to the hypothesis that it should be possible to treat tinnitus by modulating these abnormalities through the stimulation of the brain. In an initial study, Fregni et al. (2006f) targeted the left temporal cortex in 7 patients with anodal tDCS (placing the cathodal electrode supraorbital) for a single session of 3 minutes with a 1 mA intensity. They demonstrated a significant tinnitus reduction of 42% in 3 out of 7 patients. In a double-blind placebo-controlled follow-up study including 20 patients, Garin et al.
(2011) also found a significant effect of a single session of anodal tDCS of the left temporal cortex (1 mA for 20 minutes) in comparison to cathodal and sham stimulation on tinnitus loudness immediately and still one hour after stimulation. A further study investigating dose/effects revealed that single sessions of anodal tDCS of the left temporal cortex were more effective when stimulation was performed at higher stimulation intensity (2 mA) for 20 minutes (Shekhawat et al., 2013b). Using these parameters, Joos et al. (2014) found in a large series of patients with tinnitus that a single session of anodal stimulation of the left temporal cortex (39 patients), but not of the right temporal cortex (136 patients) could lead to significant tinnitus reduction. Vanneste et al. (2013a) applied tDCS at 1.5 mA for 20 minutes bilaterally on the temporal cortices (T3+T4) comparing the combination of left anode + right cathode (20 patients) versus right anode + left cathode (16 patients) and did not find any difference in the change provided by these two montages on tinnitus distress and loudness. In the same study, these authors also compared the effect of transcranial alternating current stimulation (tACS) and transcranial random noise stimulation (tRNS) applied bilaterally over the temporal cortices. A significant tinnitus reduction was observed after tRNS, but not after tDCS or tACS. Based on the hypothesis that neuroplastic changes can be promoted by tDCS, two recent studies have investigated the value of combining repeated sessions of anodal tDCS of the left temporal cortex with sound therapy (tailor-made notched music training) (Teismann et al., 2014) or hearing aids (Shekhawat et al., 2013a). However, both studies did not show any add-on effect of tDCS to the audiological treatment. Two recent studies (Forogh et al., 2016; Hyvärinen et al. 2016) were also negative. Forogh et al. (2016) reported the absence of efficacy of 5 sessions of active anodal tDCS of the left temporal cortex on tinnitus severity in 11 patients. In the study of Hyvärinen et al. (2016), two different montages were compared, either anodal tDCS of the left temporal cortex (10 patients) or a bifrontal montage with the anode on the left and the cathode on the right side (9 patients), while 11 patients received sham stimulation. These authors found no difference between active and sham groups in the effect of 10 sessions of tDCS on tinnitus severity, tDCS being applied on consecutive days as at-home treatment. Therefore, according to these replicated negative results, we can conclude that there is a level of evidence B in favour of the probable absence of efficacy of anodal tDCS of the left temporo-parietal cortex to relieve chronic tinnitus.

Other studies were designed to target the DLPFC. The rationale of DLPFC stimulation for tinnitus relief was based on neuroimaging studies showing the involvement of frontal areas in tinnitus pathophysiology (Vanneste and De Ridder, 2011; Joos et al., 2012), as well as on studies reporting clinical benefits of tDCS of the DLPFC in the treatment of major depression, which is a frequent comorbid disorder in tinnitus (Langguth et al., 2011). In a first open-labelled exploratory study (Vanneste et al., 2010), single sessions of tDCS were delivered at 1.5 mA for 20 min with the cathode over the left DLPFC (F3) and the anode over the right DLPFC (F4) (438 patients). This tDCS protocol led to a 30% suppressive effect of the tinnitus percept. Conversely, the reverse montage (left anode and right cathode) was assessed in 30 patients, but did not produce any tinnitus-suppressing effect.
Several retrospective or not-controlled single-sess ion studies of the same group further confirmed the value of bihemispheric DLPFC stimulation with left cathode and right anode for providing some tinnitus relief (Vanneste et al., 2011, 2013a,b). One study showed that bihemispheric DLPFC stimulation using tDCS could modulate tinnitus annoyance and loudness, whereas alpha-modulated tACS does not yield a similar result (Vanneste et al., 2013b). An EEG study of functional brain connectivity (coherence and phase synchronization assessment) showed that the responders to bihemispheric DLPFC tDCS differed from non-responders according to their resting brain activity in the right auditory cortex and parahippocampal area and the functional connectivity between DLPFC and the subgenual anterior cingulate cortex in particular (Vanneste et al., 2011). In an open-label study including 32 patients receiving 6 sessions of bihemispheric DLPFC tDCS with the cathode over the left DLPFC and the anode over the right DLPFC (1.5 mA, 30 min, 2 sessions per week), a small clinical effect on both tinnitus loudness and discomfort was also reported, particularly in women (Frank et al., 2012). However, in a randomized double-blind, sham-controlled study, Faber et al. (2012) found a beneficial effect on tinnitus annoyance (but not a reduction in tinnitus intensity) of anodal tDCS applied for six sessions either to the right DLPFC (7 patients) or to the left DLPFC (8 patients), with the cathode placed over the contralateral DLPFC. The effect of anodal tDCS of the DLPFC on tinnitus was irrespective of the laterality of the anode, whereas depression was preferentially modulated by left DLPFC anodal tDCS and anxiety by right DLPFC anodal tDCS. Thus, the results reported in the literature remain too preliminary to make a recommendation regarding the potential efficacy of any tDCS protocol targeted to the DLPFC for reducing either tinnitus discomfort or loudness in patients with chronic tinnitus.

Finally, we have to mention some studies using original montages, different from usually performed bipolar tDCS. First, an open-labelled study including 27 patients with chronic tinnitus (> 2 years) (Shekhawat et al., 2016) showed that a single session of tDCS with multipolar Laplacian montage centered on either the temporal cortex or the DLPFC of the left hemisphere was equally effective for suppressing tinnitus loudness and annoyance, compared to classical bipolar montage, when the stimulation was applied at 2 mA for 20 min. Second, in a large parallel-arm sham-controlled study (Pal et al., 2015), 42 patients (21 in the active group and 21 in the sham group) received 5 daily sessions of tDCS (2 mA, 20 min, 1 week) with one large anode placed over the prefrontal cortices (F3-Fz-F4) and two smaller cathodes placed bilaterally over the temporal auditory cortical areas (T3+T4). No difference in the effect of tDCS on tinnitus handicap inventory, severity scale, hospital anxiety and depression scale, and tinnitus intensity and distress was observed between the active and the sham groups from the end of the intervention up to 3 months later.

In summary, tDCS effects on tinnitus are not straightforward, depend on the methods and montages used, and have not been replicated by independent groups in large sham-controlled trials comprising multiple daily sessions. A further next step might be the evaluation of tRNS applied bilaterally over the auditory cortices, which was shown to be superior to tDCS in reducing tinnitus
distress and loudness in one single-session study (Vanneste et al., 2013a). A recently published study of tRNS of the auditory cortices in 154 patients with non-pulsatile tinnitus showed differential effects of tRNS on various tinnitus features according to stimulation frequency (Joos et al., 2015). While either low-frequency (0.1-100 Hz) or high-frequency (100-640 Hz) tRNS produced beneficial effects, the combination of both frequencies was not able to improve tinnitus. Finally, the considerable variety of design options (tDCS, tACS and tRNS), electrode montages, stimulation parameters, or frequency range concerning tACS and tRNS, opens up an unlimited amount of neuromodulation possibilities that could create a chaotic wealth of data.

Please insert TABLE 7

13. Depression

The rationale for the use of tDCS in the treatment of depressive disorders is based on the knowledge of functional and structural abnormalities in the left and right dorsolateral and ventromedial prefrontal cortex, amygdala and hippocampus in depressed patients (Campbell et al., 2004; Hamilton et al., 2008; Koenigs and Grafman, 2009). The aim of the NIBS techniques is to normalize the interhemispheric imbalance of neuronal activity between the both DLPFC areas, which was highlighted in this condition by, e.g., EEG analyses (Reid et al., 1998; Debener et al., 2000). The same rationale led to the development of rTMS therapy applied to the DLPFC, with a definite or probable evidence of antidepressant efficacy, for high-frequency rTMS of the left DLPFC or low-frequency rTMS of the right DLPFC, respectively (Lefaucheur et al., 2014).

Regarding tDCS, the current approach is to enhance neural activity in the left DLPFC with anodal stimulation and/or to reduce neural activity in the right DLPFC with cathodal stimulation (Brunoni et al., 2012). Neuroimaging and computer modeling studies of tDCS suggest that in fact, tDCS also largely affects deeper brain structures, such as amygdala, hippocampus, and subgenual cortex (Keese et al., 2011a,b; Bikson et al., 2012; Peña-Gómez et al., 2012). However, it is still unknown what changes in resting state brain networks are responsible for the antidepressant efficacy of tDCS. According to the various montages used, the respective impact of each electrode on brain network modulation also remains unclear.

A PubMed search (keywords: tDCS AND depression) identified 287 papers, including 39 original clinical studies and 988 patients, partially redundant (ancillary studies of the SELECT-tDCS trial for example). The Table 8 presents the results obtained in sham-controlled studies including at least 10 patients receiving repeated daily sessions of active anodal tDCS of the left DLPFC, according to the two main montages, with the cathode over the right orbitofrontal cortex or the right DLPFC.

**Antidepressant effects.** The antidepressant effect of anodal tDCS of the left DLPFC was first described by Fregni et al. (2006b,c). Since then, more than 10 sham-controlled RCTs were published
in this context (Fregni et al., 2006b,c; Boggio et al., 2007, 2008a; Rigonatti et al., 2008; Loo et al., 2010, 2012; Blumberger et al., 2012; Palm et al., 2012; Brunoni et al., 2013b), as well as various open-labelled studies and case reports. While RCTs investigated the therapeutic effects of tDCS in various patient samples (e.g., drug-resistant depression, unipolar versus bipolar depression) with different aims (add-on-treatment, comparison to pharmacotherapy, long-term treatment, comparison of different stimulation settings), the case reports are rather dealing with side-effects of tDCS (e.g., induction of hypomania).

Early studies used 20-min sessions of 1 mA anodal stimulation over the left DLPFC, the cathode being placed over the right supraorbital region (Fregni et al., 2006b,c). More recent trials support the use of longer sessions (30 minutes) of tDCS delivered at higher intensity (2 mA) with the cathode placed over the right DLPFC (Brunoni et al., 2013b; Valiengo et al., 2013). However, it remains to demonstrate that increasing the duration and intensity of stimulation necessarily leads to improve the therapeutic efficacy and to prolong the after-effects.

Since 2011, the antidepressant efficacy of tDCS was appraised in various meta-analyses, with the response and remission rates as primary and second outcome measures, including the calculation of effect sizes. In these works, an antidepressant response was usually defined as a 50% reduction in post-treatment scores on the Hamilton Depression Rating Scale (HDRS) or the Montgomery-Asberg Depression Rating Scale (MADRS). In the first meta-analyses, active tDCS showed no or only modest superiority compared to the sham condition (Kalu et al., 2012; Berlim et al., 2013), because of the small sample size of the included studies, a high degree of treatment resistance in at least two studies (Blumberger et al., 2012; Palm et al., 2012), or the use of tDCS as an add-on treatment to various antidepressants, leading to a ceiling effect. Furthermore, concomitant administration of mood stabilizers, benzodiazepines, or antiepileptics can influence tDCS-mediated effects on cortical excitability and theoretically might have reduced its therapeutic impact. More recent meta-analyses (Shiozawa et al., 2014; Meron et al., 2015) and an analysis of individual patient data (Brunoni et al., 2016a), including the large SELECT-TDCS trial conducted by Brunoni et al. (2013b), suggest superiority of active tDCS compared to sham treatment. In the SELECT-TDCS trial, the combination of tDCS with sertraline hydrochloride (50 mg/day) was superior to each treatment considered solely or to placebo, suggesting an additive interaction of tDCS and antidepressant pharmacotherapy. The effect of tDCS may be mediated by pharmacological modulation of serotonergic and noradrenergic neurons located in deep brain structures, although they are not directly affected by the superficial current flow generated by tDCS (Brunoni et al., 2014b). Alternatively, or additionally, serotonergic enhancement might boost the neuroplastic effects of anodal tDCS, thus resulting in synergistic effects (Nitsche et al., 2009b; Kuo et al., 2016).

Three studies have investigated the efficacy of long-term antidepressant treatment using tDCS (Dell’Osso et al., 2013; Martin et al., 2013; Valiengo et al., 2013). In a small series of 11 patients who completed a 3-month follow-up beyond a 10-day tDCS protocol, Dell’Osso et al. (2013) found that
45% of the patients were still responders at the last time point. The other studies reported a higher relapse rate when the repetition of the sessions was reduced from weekly to biweekly (Martin et al., 2013; Valiengo et al., 2013) or when the degree of treatment resistance was higher at the beginning of tDCS therapy (Valiengo et al., 2013). Other tDCS studies dealing with special issues, e.g., the influence of various biomarkers (brain-derived neurotrophic factor or cytokines), the augmentation effect with D-cycloserine administration, or the treatment of depression associated with human immunodeficiency virus infection or in the hemodialysis setting will be not reported here.

In conclusion, we have to distinguish between protocols using a cathode placed over the right orbitofrontal cortex and those using a cathode over the right DLPFC. Although the target electrode (anode) is placed over the left DLPFC in all cases, the tDCS current flow (electric field) through brain structures as well as the related network effects probably differs between the two montages. With the right orbitofrontal cathode, there is a sufficient amount of evidence (one positive class I and one positive class II studies issued from two independent groups) to make a recommendation for a level of evidence B regarding the antidepressant efficacy of anodal tDCS of the left DLPFC on the basis of at least 10 daily sessions (2 mA, 20-30 min) in medicated or drug-free patients with major depressive disorder and no drug-treatment resistance. Conversely, there is also a sufficient amount of evidence (two negative class II studies issued from two independent groups) to make a recommendation for a level of evidence B regarding the absence of efficacy of the same tDCS protocol in patients with drug-resistant depression. Concerning tDCS montage with the anode over the left DLPFC and the cathode over the right DLPFC, no recommendation can be made because of insufficient level of evidence from data published by independent groups (two positive class I-II studies issued from the same group with very few cases of drug-resistant depression and one negative class II study from another group including patients with only drug-resistant depression).

Cognitive effects. Improvement of working memory, learning, and long-term memory by tDCS has been shown in various studies performed in healthy subjects (Kuo and Nitsche, 2012). In contrast, only few therapeutic studies report a specific assessment of the cognitive impact of tDCS in patients, including patients treated for depression. Improvement of working memory in depressed patients treated by tDCS was reported by Fregni et al. (2006c) using the digit-span test and by Boggio et al. (2007) using a go/no-go task. In studies with primary neuropsychological focus, anodal tDCS of the left DLPFC was found to improve working memory in depressed subjects (Oliveira et al., 2013; Wolkenstein and Plewnia, 2013) and to modify negative emotion processing in the emotional stroop task (Brunoni et al., 2014d). One study showed, however, that bifrontal tDCS reduced implicit learning in depressed subjects (Brunoni et al., 2013c). Improvement of cognition was also mentioned in single cases of tDCS-treated patients (Palm et al., 2009; Bueno et al., 2011). Conversely, other studies reported no significant cognitive changes induced by tDCS in depressed patients (Ferrucci et al., 2009; Loo et al., 2010, 2012; Martin et al., 2011; Palm et al., 2012). Finally, one study showed that
the combination of cognitive control training (a new type of cognitive therapy) and tDCS has an augmenting effect on the improvement of depression (Segrave et al., 2014).

Concerning the impact of tDCS on cognitive symptoms of patients with major depression, a beneficial effect was reported in four class II studies, whereas one class I study and three class II studies were negative. In particular, two positive and one negative study emerged from the same SELECT-TDCS trial. Therefore, current data remain conflicting and no recommendation can be made for the use of tDCS of the DLPFC to improve cognitive symptoms in depressed patients. This conclusion is shared by an article recently published on this topic (Brunoni et al., 2016b).

**Please insert TABLE 8**

### 14. Schizophrenia

Despite advances in psychopharmacology, the majority of treated patients with schizophrenia retain disabling symptoms. The most frequent drug-resistant symptoms are auditory verbal hallucinations (AVH) and various negative symptoms (e.g., avolition, alogia, or emotional withdrawal). Neuroimaging and neurophysiological studies have highlighted that these refractory symptoms may be linked to a fronto-temporal dysconnectivity. Negative symptoms and AVH have been related to a reduced brain activity in the right and left DLPFCs, while hyperactivity in the left temporo-parietal region is especially observed in patients with AVH. Using the concept of excitatory anodal stimulation versus inhibitory cathodal stimulation, it was hypothesized that anodal tDCS of the left DLPFC (hypoactive) combined with cathodal tDCS of the left temporo-parietal junction (hyperactive) should alleviate negative symptoms and AVH in patients with schizophrenia (Brunelin et al., 2012b; Mondino et al., 2014). In the same way, bilateral tDCS delivered to both prefrontal regions (left anode plus right cathode) should alleviate negative symptoms.

A PubMed search (keywords: tDCS AND schizophrenia) identified 116 papers, including 38 original clinical studies and 384 patients. Among these 37 papers, there were 19 case reports and 19 clinical studies with different tDCS protocols applied on samples of patients with various symptoms and group sizes. In fact, for the present study, we retained only 7 sham-controlled RCTs including at least 10 patients who received repeated daily sessions of active tDCS with the anode placed over the left DLPFC (between F3 and FP1 according to the International 10-20 system of EEG electrode placement) (Brunelin et al., 2012a; Fitzgerald et al., 2014; Mondino et al., 2015, 2016; Smith et al., 2015; Fröhlich et al., 2016; Palm et al., 2016b). In two of these studies, the cathode was placed on the right supraorbital region (Smith et al., 2015; Palm et al., 2016b) and they will be detailed later. In the remaining 5 studies, the cathode was placed over the left temporo-parietal junction (mid-point between T3 and P3) and the effects of this bipolar montage over a large fronto-temporo-parietal region of the left hemisphere were assessed on various positive or negative symptoms of schizophrenia (Table 9).
Among these 5 studies, one double-blind sham-controlled RCT showed a significant reduction of AVHs following active tDCS compared to sham condition (Brunelin et al., 2012a). The protocol consisted of 10 daily sessions of tDCS (2 mA, 20 min) over two consecutive weeks. Clinical improvement was significant on persistent AVHs, but also on both negative and positive dimensions, assessed on the total Positive And Negative Symptoms Scale (PANSS) as a secondary outcome. The same group showed in partly overlapping samples that the reduction in AVHs following this type of tDCS protocol correlated with an improvement in source-monitoring performance (Mondino et al., 2015) and with a reduction of functional connectivity between the left temporo-parietal junction and anterior insula (Mondino et al., 2016). In contrast, little is known on the ability of this approach to improve cognitive functions, such as spatial working memory or speech processing, which are altered in patients with schizophrenia. A double-blind sham-controlled feasibility study confirmed the safety of this intervention in a series of 12 young patients with childhood-onset schizophrenia (Mattai et al., 2011). However, two replication studies, including one comprising two small RCTs (unilateral and bilateral tDCS), and published by independent groups failed to demonstrate any clinically relevant effect of active vs. sham tDCS using this montage on AVHs and negative symptoms at the end of the intervention based on 15 tDCS sessions over three weeks (Fitzgerald et al., 2014) or 5 tDCS sessions over one week (Fröhlich et al., 2016). However, these studies suffered from various weaknesses, including small sample size, leaving room for further replication studies or large multicentric RCTs using multiple-session tDCS treatment in schizophrenic patients with the anode over the left DLPFC and the cathode over the left temporo-parietal cortex. Meanwhile, we cannot make any recommendation about the efficacy of this montage to relieve schizophrenia symptoms, either positive or negative.

Other studies used a different electrode montage, with the anode still placed over the left DLPFC, but the cathode placed over the right supraorbital region. Using this montage, some beneficial results were reported in patient cases concerning the relief of negative symptoms of schizophrenia (Palm et al., 2013b) or catatonia (Shiozawa et al., 2013). In two controlled studies using this montage (Smith et al., 2015; Neniow et al., 2016), the cognitive impairment and deficits associated with schizophrenia were assessed, while schizophrenia symptoms were secondary outcomes or not assessed. These studies showed the positive effects on cognition of repeated sessions of anodal tDCS of the left DLPFC performed prior or during cognitive training. In the first study, with a parallel-arm design, 29 patients (14 in the active group and 15 in the sham group) received 5 sessions of anodal tDCS (2 mA, 20 min) of the left DLPFC (with right supraorbital cathode) over one week (Smith et al., 2015). A significant cognitive improvement was observed at the end of the intervention after active vs. sham tDCS in composite, working memory and attention-vigilance scores of the MATRICS Consensus Cognitive Battery. However, there were no differences between the two tDCS conditions regarding the PANSS scores (including AVH and negative symptoms) and cigarette smoking or craving. A single-blind, sham-controlled, proof-of-concept study also addressed cognitive issues in 10
schizophrenic patients receiving 28 sessions (two sessions each week) of tDCS combined with cognitive training (Nienow et al., 2016). The 6 patients who received active stimulation with the anode over the left DLPFC and the cathode over the contralateral supraorbital area showed significant more improvement on word or picture 2-back tasks compared to the 4 patients who received sham stimulation. However, schizophrenia symptoms were not assessed. Conversely, the most recent study using this montage had the Scale for the Assessment of Negative Symptoms (SANS) as the primary outcome measure (Palm et al., 2016b). This double-blind sham-controlled study with parallel-arm design (10 patients per arm) showed a significant reduction of 36% of negative symptoms on SANS after 10 active stimulations compared to a reduction of 0.7% in the sham group. As secondary outcome, PANSS sum scores also decreased significantly more after active (−23.4%) than sham stimulation (−2.2%). However, the results of the two sham-controlled studies using this montage (Smith et al., 2015; Palm et al., 2016b) being conflicting (Table 9), no recommendation can be made for the use of anodal tDCS of the left DLPFC with a contralateral orbitofrontal cathode in schizophrenia, even regarding cognitive deficits.

Finally, some results were reported in this domain, still with the anode over the left DLPFC but with other cathode placements. For example, with the cathode placed over the right DLFPC (bilateral DLPFC stimulation), a small RCT, including 15 patients and specifically addressing the treatment of negative symptoms, showed a significant decrease in PANSS total score and negative subscale after active stimulation compared to sham condition (Gomes et al., 2015). Using a cathode placed over the right deltoid muscle, an open-label study of 9 patients with negative symptoms also found an improvement of 24% in the PANSS negative subscale after anodal tDCS of the left DLPFC (Kurimori et al., 2015). The results obtained with these montages, non-replicated yet, cannot be considered for making recommendation.

Please insert TABLE 9

15. Substance abuse, addiction and craving

Addiction to substances such as alcohol, drugs, nicotine, or food, is a major health issue, because of the difficulty to achieve a permanent cure with a high rate of relapses, despite detoxification and pharmacological or psychological interventions (Fant et al., 2009; Heinz et al., 2009). The rationale of using tDCS as a treatment for substance addiction and craving is that the DLPFC, which plays a major role in top-down inhibitory control mechanisms and reward mechanisms, was claimed to be dysfunctional in these disorders (Goldstein and Volkow, 2002; Wilson et al., 2004).

A PubMed search (keywords: tDCS AND addiction OR craving) identified 80 papers, including 32 original clinical studies and 583 subjects. No safety problems emerged in these studies. There were 19 studies including at least 10 patients, but mostly based on single tDCS sessions with
immediate or short-term assessment. In fact, we only identified 4 sham-controlled studies with at least 10 patients who received repeated daily sessions of the same type of active tDCS, i.e. a bihemispheric stimulation of the DLPFC with the anode on the right (F4) and the cathode on the left hemisphere (F3) (Boggio et al., 2009b; Fecteau et al., 2014; Klauss et al., 2014; Batista et al., 2015) (Table 10). The analyzed results cover patients addicted to alcohol, cocaine, crack, or smoking. All these studies were positive regarding the effect of active tDCS, either in the resulting quality of life (Klauss et al., 2014; Batista et al., 2015) or in craving scores (Boggio et al., 2009b; Fecteau et al., 2014; Batista et al., 2015). According to these four positive class II studies issued from two independent teams, we can make a recommendation for a level of evidence B regarding the efficacy of bihemispheric tDCS of the DLPFC (right anode + left cathode) to reduce craving in patients with various types of addiction. However, this statement must be tempered by the fact that studies concerning alcohol abuse, crack-cocaine abuse, and smoking were pooled together, with results obtained on different outcome variables. Nevertheless we want to keep this statement, since a common mechanism of action certainly underlies all the results obtained with this given montage in patients who all have an addiction. This conclusion is reinforced by a recent neurophysiological and neuroimaging study showing that abstinence provided by bihemispheric tDCS of the DLPFC using this specific montage (right anode + left cathode) could involve functional changes in the ventral medial prefrontal cortex in both alcoholics and crack-cocaine users (Nakamura-Palacios et al., 2016).

It is interesting to note that this montage is similar to the one proposed in the treatment of tinnitus, but opposed to the one proposed in the treatment of depression (anodal tDCS of the left DLPFC with right DLPFC cathode) (see previous chapters). In fact, as aforementioned, Faber et al. (2012) found a beneficial effect on tinnitus regardless of the direction of the montage between the two hemispheres, but tinnitus-associated depression was preferentially relieved by placing the anode over the left DLPFC, whereas anxiety was preferentially relieved by placing the anode over the right DLPFC.

Actually, in alcoholics, one study showed a reduction of craving for visual alcohol cues after a single session of active bihemispheric tDCS of the DLPFC, similarly for a right anode + left cathode montage and a left anode + right cathode montage (Boggio et al., 2008b). Conversely, the same team showed that only the right anode + left cathode montage was significantly associated with a reduction of craving for marijuana, while both montages increased the propensity for risk-taking in chronic marijuana users (Boggio et al., 2010). Other working groups interestingly found that only the right anode + left cathode montage was specifically able to reduce the risk-taking behavior in 18 dependent cocaine users (Gorini et al., 2014) and 18 cigarette smokers (Pripfl et al., 2013). This right anode + left cathode montage, applied in single or multiple tDCS sessions (2 mA, 20 min, 5 days for one week) was found to positively impact cognitive processing related to crack-related visual cues in 13 crack-cocaine users (Conti et al., 2014). Pripfl and Lamm (2015) also showed that a single session of bihemispheric tDCS of the DLPFC with the right anode + left cathode montage (and not the reverse
montage) could reduce negative affect in emotion appraisal but not modulate nicotine craving-cue appraisal in 20 smokers. Finally, Kekic et al. (2014) reported that a single tDCS session with the right anode + left cathode montage over the DLPFC improved craving scores for sweet but not savoury foods in 17 patients with food craving. The participants that exhibited more reflective choice behaviour were more susceptible to the anti-craving effects of tDCS than those that displayed more impulsive choice behaviour. However, no difference in food consumption was observed after active vs. sham tDCS. Conversely, one week of daily anodal tDCS delivered to the right DLPFC (but with the cathode placed in the left supraorbital region) was found to reduce overall caloric intake by 14% in comparison with sham stimulation in 14 subjects, which however do not meet the criteria of eating disorder (Jauch-Chara et al., 2014).

Other studies used a tDCS montage with the anode over the left DLPFC and a cathode over the contralateral supraorbital or shoulder region. Using this montage, a transient improvement on clinical and electrophysiological features of craving was observed in alcoholics after a single active tDCS session (Nakamura-Palacios et al., 2012; den Uyl et al., 2015) or a series of 5 sessions (once a week for five consecutive weeks) (da Silva et al., 2013). In smokers, anodal stimulation of the left DLPFC with cathode placed over the contralateral supraorbital region was found to reduce the negative affect correlated with nicotine dependence without any change on cigarette craving in one single-session study (Xu et al., 2013) or to improve the ability to resist smoking in another single-session study (Falcone et al., 2016). Obviously, these results are still too preliminary to make any recommendation for the use of anodal tDCS of the left DLPFC with the cathode over the right orbitofrontal region in craving. Finally, the reverse tDCS montage, with the anode over the right DLPFC and the cathode over the left supraorbital region, was recently shown to be able to reduce food craving (Ljubisavljevic et al., 2016). A single session could produce immediate effects, while the clinical improvement could last for a month after a 5-day protocol. These results remain to be replicated.

Please insert TABLE 10

16. Other psychiatric disorders

Regarding clinical applications of tDCS in psychiatric disorders other than depression, schizophrenia and addiction, there are only very limited data available in the literature. A PubMed search (keywords: tDCS AND obsessive compulsive disorder) identified 17 papers, including 5 original clinical studies and 12 patients. There were 4 single case reports and only one open-label study performed on 8 patients with the cathode placed over the left orbitofrontal cortex and the anode placed over the right cerebellum (Bation et al., 2016). A PubMed search (keywords: tDCS AND post-traumatic stress disorder) identified 5 papers, including only one pilot study of 4 patients (Saunders et
A PubMed search (keywords: tDCS AND autism) identified 36 papers, including 6 original clinical studies and 64 patients. Among these 6 papers, there were 2 single case reports and 2 RCTs issued from the same team assessing the effect of either a 5-day protocol on clinical scores or a single session on EEG parameters in a series of 20 patients (Amatachaya et al., 2014, 2015). The montage used was an anode placed over the left DLPFC (F3) with a cathode on the right shoulder. Finally, a PubMed search (keywords: tDCS AND attention-deficit hyperactivity disorder) identified 24 papers, including 2 original clinical studies (single-session studies) and 80 patients. One study (60 patients) did not find significant differences between active and sham tDCS regarding the effect of bihemispheric tDCS of the DLFPC (left anode + right cathode) on behavioral performance in go/no-go tasks (Cosmo et al., 2015). Conversely, the other study (20 patients) showed that anodal tDCS of the left DLPFC (with right supraorbital cathode) increased the proportion of correct responses in go/no-go tasks, whereas cathodal tDCS of the left DLPFC increased the inhibition accuracy (Soltaninejad et al., 2016). Obviously, reported results are too preliminary to make any recommendation for the use of tDCS in all these psychiatric conditions.

17. At-home do-it-yourself DCS and neural enhancement

One of the main advantages of tDCS as a therapeutic device is to be a low-cost, small-size, and patient-wearable equipment. Also, it allows distribution of machines for home use, which is not the case for rTMS. Although therapeutic effects may last beyond the time of stimulation when using NIBS techniques because of neural plasticity processes, the treatment of a chronic disease always requires repeated sessions in a maintenance protocol. Considering rTMS, this is resulting in multiple trips and visits for the patient to be stimulated, generating fatigue and costs, and reducing compliance. Considering tDCS, at-home application is feasible and potentially more comfortable and less tiring for the patient. At-home tDCS trials have been already reported as case reports for the treatment of AVH (Andrade, 2013) or chronic myofascial pain (Pérez-Borrego et al., 2014), series of patients with trigeminal neuralgia (Hagenacker et al., 2014) or tinnitus (Hyvärinen et al., 2016), or are currently under investigation in patients with chronic pain (O'Neill et al., 2015) or MS (Kasschau et al., 2015). One group also used tDCS as an add-on intervention combined with physical therapy to promote motor recovery in patients impaired by intracerebral hemorrhage (Mortensen et al., 2016). Recommendations have even been recently proposed for a safe use of remotely-supervised at-home tDCS (Charvet et al., 2015), highlighting training of the user or caregiver, medical supervision, monitoring of compliance, and assessment of the clinical benefit or side-effects. However, an uncontrolled domiciliary utilization of tDCS devices exposes the patient to potential adverse events caused by misuse or overuse, e.g., skin burns (Wang et al., 2015) or unnecessary or dangerous repetition of the sessions.
Because of the wide diffusion of tDCS or tDCS-like machines that can be bought freely on the internet, it will be difficult to restrict the use of these machines and ensure correct application. This also refers to the topic of “neuroenhancement”, which is however beyond the scope of this present work, as it relates to improving the capacities of normal subjects and not to treat pathological conditions. In this context of free access to technology, neuroenhancement by tDCS is the subject of many ethical and regulatory debates (Hamilton et al., 2011; Cohen Kadosh et al., 2012; Heinrichs, 2012; Santarnecchi et al., 2013; Cabrera et al., 2014; Fitz and Rainer, 2015; Maslen et al., 2015; Wexler, 2015) and scientific controversies. In fact, theoretically, tDCS may have the capacity to “enhance” attention, learning, endurance, motor execution, memory, or other higher-order processes such as decision-making, risk-taking, or problem-solving (Levasseur-Moreau et al., 2013; Brem et al., 2014; Coffman et al., 2014). However, tDCS experts’ opinions are not unequivocal on this topic, as shown by two recent surveys (Shirota et al., 2014; Rigall et al., 2015). In fact, modifying neural activity by NIBS does not always go in the expected direction in a “normal” human brain. For example, one study showed that the improvement of cognitive performance by testing repetition was reduced in healthy subjects after a single session of active tDCS over the DLPFC (bilateral or unilateral anodal stimulation with the cathode placed at Cz) compared to the sham procedure (Sellers et al., 2015). Specifically, the reduction in test-retest improvement was observed on the Full Scale Intelligence Quotient and the Perceptual Reasoning Index of the Wechsler Adult Intelligence Scale. Another study showed that the accuracy performance in a working memory task was significantly decreased during and after a single session of tDCS over the DLPFC using a commercial tDCS machine that is freely available on the internet as a cognitive enhancer (Steenbergen et al., 2015). Therefore, there is a need for maintaining a level of technical and scientific excellence and considering regulatory issues to not discredit this therapeutic perspective. In this context, the International Federation of Clinical Neurophysiology (IFCN) recently warns against the use of DIY devices and methods of NIBS unless they have shown both efficacy and safety (see recommendation in the following document: http://www.ifcn.info/uploadfiles/documents/2015/Using_tES_devices_as_DIY_FINAL_13Dec15.pdf).

18. Perspectives of targets other than cortical (cerebellum and spinal cord)

Although this work is intended to be a guideline on the indications of tDCS applied to the cerebral cortex, readers should be informed that research currently develops to explore the possibility of using transcutaneous DC stimulation on other neural targets, such as cerebellum and spinal cord, in order to promote functional neural changes (Priori et al., 2014). This issue will be briefly addressed.

Cerebellar tDCS. Because the cerebellum plays a major role in several brain functions, cerebellar stimulation could be a preferential way of therapeutic intervention in many pathological conditions. Weak DC delivered transcutaneously in humans over the cerebellum for minutes elicits prolonged changes in neurophysiological and behavioural responses related to cerebellar functions.
Theoretical considerations and modelling studies suggest that the electric field generated by tDCS can reach the cerebellum when using an appropriate montage (Parazzini et al., 2014b). The induced effects arise from functional changes in the stimulated cerebellum, although transynaptic or antidromic changes may also occur in connected cerebral or brainstem structures.

There is already reported evidence that cerebellar tDCS can induce neurophysiological, functional, and clinical effects related to DC-induced neuroplasticity and neurotransmitter changes (Ferrucci et al., 2015; Grimaldi et al., 2016). For example, cerebellar tDCS was found to influence cerebello-thalamo-cortical inhibitory projections (Galea et al., 2009), human associative plasticity assessed by the paired-associative stimulation paradigm (Hamada et al., 2012), somatosensory mismatch negativity (Chen et al., 2014) conditioned eyelink responses (Zuchowski et al., 2014), gait adaptation (Jayaram et al., 2012), motor learning (Galea et al., 2012; Dutta et al., 2014; Hardwick and Celnik, 2014; Herzfeld et al., 2014), and cognitive functions, such as memory and emotion (Ferrucci et al. 2008b, 2012, 2013; Boehringer et al., 2012; Pope and Miall, 2012; Macher et al., 2014). These results have been obtained in healthy humans and could have clinical interest, especially in patients with disorders involving cerebellar dysfunction such as ataxia, Parkinson's disease, autism and schizophrenia (Yeganeh-Doost et al., 2011; Massaquoi, 2012; Wu and Hallett, 2013).

There are only preliminary data of the application of cerebellar tDCS in patients (Grimaldi and Manto, 2013; Bradnam et al., 2014, 2015; Gironella et al., 2014; Grimaldi et al., 2014b; Minichino et al., 2014, 2015; Sadnicka et al., 2014; Benussi et al., 2015; Bation et al., 2016; Ferrucci et al., 2016). Much work remains to be done to design further therapeutic studies using cerebellar tDCS, according to the parameters of stimulation (electrode montage, polarity), the possible mechanisms of action, and the underlying pathological conditions and interactions with ongoing drug treatments.

**Transcutaneous spinal direct current stimulation.** Invasive high-frequency epidural electrical spinal cord stimulation (SCS) has been used for more than 30 years to treat a variety of pain syndromes (Grabow et al., 2003; Mailis-Gagnon et al., 2004; Frey et al., 2009). Involving totally different mechanisms of action, a noninvasive approach of spinal cord neuromodulation was recently developed, using transcutaneous DC stimulation in humans (Cogiamanian et al., 2008, 2012).

The application of anodal DC at the thoracic level (2.5 mA for 15 min with the cathode placed at the right shoulder) selectively reduced the amplitude of the cervicomедullary P30 component of the somatosensory evoked potentials to posterior tibial nerve stimulation (Cogiamanian et al. 2008) as well as laser evoked potentials after foot stimulation (Truini et al. 2011) or nociceptive withdrawal reflex (Cogiamanian et al., 2011; Perrotta et al., 2016). These results provide evidence that transcutaneous spinal direct current stimulation (tSICS) is able to interfere with spinal cord conduction properties and to modulate conduction in the lemniscal and spino-thalamic pathways. At present, there is also increasing evidence that tSICS is able to induce persisting changes in properties of spinal sensory or motor neurons (Winkler et al., 2010; Lim and Shin 2011; Lamy et al., 2012; Hubli
et al., 2013; Niérat et al., 2014). The resulting excitability changes induced by tsDCS may also extent to corticospinal tracts or even intracortical circuits (Bocci et al., 2015a,b,c).

To date, the precise mechanisms underlying these neuromodulatory effects remain speculative, although one modelling study improves the understanding of the spatial distribution of the current density generated by tsDCS (Parazzini et al., 2014a). Nevertheless the possibility of influencing conduction along the ascending spinal pathway as well as spinal circuitries is especially interesting for clinical purposes including the treatment of chronic pain syndrome, e.g., restless leg syndrome (Heide et al., 2014) and the neurorehabilitation of motor disorders, especially spasticity or poststroke gait impairment (Picelli et al., 2015).

19. Perspectives of treatment by transcranial electrical stimulation methods other than tDCS (tACS, tRNS)

Although the present work addresses tDCS results, we should also mention that future developments with potential therapeutic application might regard other non-invasive neuromodulation techniques using low-intensity transcranial electrical stimulation. We briefly present these other techniques, namely tACS and tRNS, which are for now the subject of relatively few clinical studies. Taking into account all physical possibilities, an indefinite number of transcranial electrical stimulation protocols exists. Rather arbitrarily determined protocols have been tried over the last century, mainly deriving from the Franco-Russian experience of “electrosleep” or “electroanesthesia” (Guleyupoglu et al., 2013). Most of the complex historical stimulation protocols are proprietary commercial “blends” and will probably not survive the test of time against physically simpler and better evaluated protocols such as tDCS or sinusoidal tACS (Paulus, 2011). In case of tDCS, first clinical descriptions date as early as 1801, one year after the invention of the voltaic pile. An earlier attempt of modulating slow EEG oscillations by tACS was performed during sleep, but tACS was applied with tDCS overlay, leaving until today unclear whether the DC or the AC component provided the resulting effect (Marshall et al., 2006). Then, the first investigation with pure sinusoidal motor cortex tACS delivered at 10 Hz produced only small aftereffects on implicit motor learning (Antal et al., 2008). This was probably due to the low intensity of stimulation (0.4 mA), which was chosen to avoid retinal phosphenes via the frontal return electrode. Since a couple of years, tACS was applied in disease context, first in movement disorders (Angelakis et al., 2013) and tinnitus (Vanneste et al., 2013a,b), and more recently to modulate tremor by targeting the cerebellum (Mehta et al., 2014).

Two different goals have to be differentiated when looking at tACS effects, first regarding entrainment with ongoing brain rhythm and second regarding induction of aftereffects. In the context of the present review, the latter seems to be the most interesting since this would avoid wearing the stimulation electrode all day for therapeutic purposes. The problem of the necessity for continuous stimulation was pointed out by a closed-loop tACS approach using specified phase alignments to
suppress parkinsonian tremor by about 50% (Brittain et al., 2013). This is also a requirement for the application of alternating fields at very high intensity and frequency for therapeutic purpose in a disease justifying substantial efforts for treatment, namely the treatment of glioblastoma (Swanson et al., 2016). The upper frequency limit of tACS has not yet been determined and it touches e.g. the question if mobile phone emission can influence brain function. In fact, continuous tACS at 200 kHz could selectively destroy dividing cells by targeting the inhomogeneous fields present at the bridge separating the daughter cells that interfere with spindle tubulin orientation and induce dielectrophoresis (Kirson et al., 2007). Conversely, the role of tACS frequencies in the low kHz range remains to be investigated. A first study showed that 2 and 5 kHz tACS protocols were particularly efficacious in inducing a sustained increase in MEP amplitude up to 30-60 min after the intervention (Chaieb et al., 2011). Within this frequency, tACS appears too fast to entrain brain circuit oscillations, and therefore the stimulation probably directly interferes with cortical excitability via neuronal membrane activation changes.

A protocol of 140 Hz tACS delivered at 1 mA for 10 minutes can produce aftereffects comparable in duration to those induced by anodal tDCS (Moliadze et al., 2010). This frequency complies with the so-called “ripple frequency”, which has a function in encoding memory in the hippocampus. Interestingly it is about the same frequency which was empirically found to be optimally suited for DBS therapeutical effects in movement disorders (130 Hz). In the context of tACS, stimulation intensity plays a critical role to promote either excitation or inhibition of the neural circuits (Moliadze et al., 2012) and not electrode polarity as for tDCS. A deeper understanding is requested to determine how electrode size, shape, or montage, as well as stimulation frequency and intensity (Moliadze et al., 2012) and the initial state of the brain and its susceptibility to resonance effects (Feurra et al., 2013) may interfere in a complex manner in the emergence of tACS effects beyond the simple process of oscillation entrainment or spiking resonance (Reato et al., 2013; Schmidt et al., 2014).

Although motor cortex excitability parameters assessed by TMS, e.g., motor threshold or MEP size measurement, provides widely used biomarkers, their value in predicting behavioural changes induced by tACS remains highly speculative. For example, 140 Hz tACS was very efficient in influencing MEP size, whereas implicit motor learning was rather facilitated by 250 Hz tACS (Moliadze et al., 2010). Frequency-specific effects can be circumvented by performing tRNS, which was introduced by Terney et al. (2008) as a novel method of transcranial electrical stimulation. However, a random noise spectrum using frequencies ranging from 100 to 640 Hz led essentially to the same results as 140 Hz tACS or anodal tDCS, i.e. a consistent increase in motor cortex excitability (Terney et al., 2008). In fact, tRNS may provide larger MEP size increase and anodal tDCS longer MEP size increase compared to sham, suggesting different underlying mechanisms of action (Moliadze et al., 2014). As tACS, tRNS was also applied in disease context since a couple of years, to provide “therapeutic” improvement, such as the relief of neuropathic pain (Alm and Dreimanis, 2013;
Palm et al., 2016a), psychiatric symptoms (Palm et al., 2013a; Haesebaert et al., 2014), or tinnitus (Vanneste et al., 2013a,b; Claes et al., 2014; Joos et al., 2015). These reported clinical effects should be specifically assessed, but largely remain preliminary before being replicated by independent groups in sham-controlled studies.

20. Summary of recommendations

This work presents for the first time a comprehensive evidence-based analysis of the reported clinical efficacy of various tDCS montages that could lead to therapeutic applications in the neurological, otorhinolaryngological, and psychiatric domains. According to this synthesis, there is a sufficient level of evidence to make recommendations for the efficacy of specific tDCS montages in several clinical indications, as summarized in Table 11.

To date, a Level A recommendation has not been achieved so far for any clinical indication. In contrast, a Level B recommendation (probable efficacy) is conferred for the clinical effect of: (i) anodal tDCS of the left M1 (defined as C3 location in the International 10-20 system of EEG electrode placement) with the cathode over the right orbitofrontal region in fibromyalgia; (ii) anodal tDCS of the left DLPFC (F3) with the cathode over the right orbitofrontal region in non-drug-resistant major depressive episodes; (iii) anodal tDCS of the right DLPFC with the cathode over the left DLPFC in addiction/craving, although this statement covers different clinical conditions and outcome variables. A Level C recommendation (possible efficacy) is conferred for the clinical effect of anodal tDCS of the motor cortex of the left hemisphere or contralateral to pain side with the cathode over the right orbitofrontal region in patients with chronic neuropathic pain at the lower limbs secondary to traumatic or medical spinal cord lesion.

Conversely, a Level B recommendation (probable inefficacy) is conferred for the absence of clinical effect of: (i) anodal tDCS of the left temporal cortex (between C3 and T3/T5) with the cathode over the right orbitofrontal region in chronic tinnitus; (ii) anodal tDCS of the left DLPFC (F3) with the cathode over the right orbitofrontal region in drug-resistant major depression. Finally, in clinical conditions where no recommendation was proposed, it must be underlined that the absence of evidence should not be taken as evidence for the absence of effect.

Further controlled studies in all potential therapeutical indications are obviously needed to extend and confirm the present recommendations. The term “recommendation” used in this work should not be misinterpreted. Our objective was to “recommend” a type of tDCS protocol in a given clinical situation according to the level of evidence of its “real” efficacy, comparing active vs. sham condition. Our methodology was not that of a meta-analysis using all data published in a statistical way, but was based on the existence of results replicated by independent teams on a representative sample of patients who received multiple sessions of active stimulation. For example, the beneficial results provided by a given tDCS protocol in a specific indication but only reported by one team and
not replicated by independent teams did not lead to a recommendation, whereas such results might have a significant weight in a meta-analysis. In addition, a high level of evidence of efficacy (comparing active vs. sham condition) should not open the door to clinical use in daily routine practice without any limitations. The first of these limitations is to precisely design the tDCS protocol to apply, especially in terms of session duration and repetition or combined therapy. Our work was intended to present evidence-based guidelines, not practical guidelines to codify the daily use of tDCS therapy. The second limitation is the fact that a “significant” effect of active stimulation compared to sham control may be statistically relevant but not clinically meaningful in daily living or in terms of quality of life. Therefore, it remains to be determined how to optimize tDCS protocols and techniques to give them “therapeutic relevance” in routine clinical practice. The safety profile of tDCS is very high, as far as explored, with regard to the currently applied protocols (Bikson et al., 2016). Adverse effects are limited to mild headache or local sensory discomfort (Poreisz et al., 2007; Brunoni et al., 2011; McFadden et al., 2011; Kessler et al., 2012). In practice, the main issue is to avoid local skin burns by limiting excessive current density according to electrode size and shape, keeping a good homogeneous contact between the electrodes and the skin using gel, cream, or appropriately large, wet electrodes (Loo et al., 2011; Palm et al., 2014b). In this context, technical guidelines have been recently proposed by a group of experts to ensure a proper use of tDCS protocols (Woods et al., 2016).

Our work has other limitations. First, our literature search strategy was based on only one database (PubMed), but this is one of the main international databases of references on life sciences and biomedical topics. In addition, the research was conducted by experts in the field. Therefore, it is unlikely that original published data of importance can miss in this work. More importantly, we cannot rule out the impact of publication bias on our analysis, which corresponds to the fact that trials with statistically significant beneficial results are most likely to be submitted and published than trials with "negative" or non-significant results. This is a well known problem for any conclusion reported in literature reviews or meta-analyses based only on published studies, which should be considered with caution, even when analysis methods seem very stringent (Easterbrook et al., 1991). Finally, it is important to underline that our analysis was based on arbitrary criteria regarding the number of patients receiving active treatment needed to be included (≥25 in a class I study and ≥10 in a class II study), which could be considered relatively small to people outside of this field of research. The results of the analyses and therefore the recommendations would have changed if the number of patients receiving active treatment needed to be included was set at a higher value (e.g., ≥15-20 patients in the "active" arm rather than ≥10 patients), mostly because few studies would exceed this cutoff. However, the main interest and justification of the method used in this work are to be identical to the previously published recommendations for therapeutic use of rTMS (Lefaucheur et al., 2014) and thus allowing comparisons.

We must be aware of the existence of current technical developments, focussing on new designs of electrode montages, shapes or sizes, resulting in new “geometries” of electric field
distribution within the brain (Saturnino et al., 2015), or on combined approaches with neuroimaging (e.g., fiber tracking) and high-resolution EEG to improve targeting. Such developments, as well as a better characterization of stimulation intensity, duration, and session repetition requirements, may help to reduce the large interindividual variability in tDCS efficacy that currently renders the average clinical responses rather modest. The use of tDCS should also be considered as an adjunctive therapy in combination with medication or non-pharmacological therapy, with the aim of improving or accelerating the efficacy of these treatments. However, various applications of tDCS in combined protocols with concomitant interventions showed no additional impact of tDCS, possibly due to a ceiling effect, as shown for the combination of tDCS and robot-assisted training in motor rehabilitation of stroke patients (Geroin et al., 2011; Hesse et al., 2011; Triccas et al., 2015). Furthermore, different drug treatments may even reduce tDCS efficacy, leading to a worse outcome compared to unmedicated patients, as shown for the use of benzodiazepines in depressed patients treated by 5 tDCS sessions, with the anode over the left DLPFC and the cathode over the right DLPFC (Brunoni et al., 2013a).

To conclude, there are pros and cons for the different NIBS techniques, such as tDCS and rTMS, applicable in clinical conditions. The respective place of these techniques remains to be determined in the available armamentarium and the decision tree for the treatment of neuropsychiatric diseases. However, it is likely that the indications of NIBS techniques will increasingly develop in routine clinical practice in the future, mostly due to their excellent ratio between benefit and risk. These guidelines pave the way for the therapeutic application of tDCS by showing already acquired evidence of efficacy of this technique in the current literature.

Conflict of interest

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References
Andrade C. Once- to twice-daily, 3-year domiciliary maintenance transcranial direct current stimulation for severe, disabling, clozapine-refractory continuous auditory hallucinations in schizophrenia. J ECT 2013;29:239-42.


Bindman LJ, Lippold OCJ, Redfearn JWT. The action of brief polarizing currents on the cerebral cortex of the rat (1) during current flow and (2) in the production of long-lasting after-effects. J Physiol (London) 1964;172:369-82.


Brunoni AR, Júnior RF, Kemp AH, Lotufo PA, Benseñor IM, Fregni F. Differential improvement in depressive symptoms for tDCS alone and combined with pharmacotherapy: an exploratory analysis from the sertraline vs. electrical current therapy for treating depression clinical study. Int J Neuropsychopharmacol 2014b;17:53-61.


Faria P, Hallett M, Miranda PC. A finite element analysis of the effect of electrode area and inter-electrode distance on the spatial distribution of the current density in tDCS. J Neural Eng 2011;8:066017.


Fröhlich F, Burrello TN, Mellin JM, Cordle AL, Lustenberger CM, Gilmore JH, Jarskog LF. Exploratory study of once-daily transcranial direct current stimulation (tDCS) as a treatment for auditory hallucinations in schizophrenia. Eur Psychiatry 2016;33:54-60.


Grüner U, Eggert C, Ameli M, Sarfeld AS, Fink GR, Nowak DA. 1 Hz rTMS preconditioned by tDCS over the primary motor cortex in Parkinson's disease: effects on bradykinesia of arm and hand. J Neural Transm 2010;117:207-16.


Kiran S. What is the nature of poststroke language recovery and reorganization? ISRN Neurol 2012;2012:786872.


Saturnino GB, Antunes A, Thielischer A. On the importance of electrode parameters for shaping electric field patterns generated by tDCS. Neuroimage 2015;120:25-35.


Shekhawat GS, Stinear CM, Searchfield GD. Transcranial direct current stimulation intensity and duration effects on tinnitus suppression. Neurorehabil Neural Repair 2013b;27:164-72.


Vanneste S, Fregni F, De Ridder D. Head-to-Head Comparison of Transcranial Random Noise Stimulation, Transcranial AC Stimulation, and Transcranial DC Stimulation for Tinnitus. Front Psychiatry 2013a;4:158.


Zuchowski ML, Timmann D, Gerwig M. Acquisition of conditioned eyeblink responses is modulated by cerebellar tDCS. Brain Stimul 2014;7:525-31.
Table 1. Repeated tDCS session protocols in chronic pain conditions

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
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<tr>
<td><strong>Chronic neuropathic pain of the lower limbs due to spinal cord lesion</strong></td>
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<td>Fregni et al., 2006a</td>
<td>17 patients with traumatic spinal cord injury at various anatomical levels (11 active, 6 sham)</td>
<td>Anode: M1 (C3/C4) of the dominant hemisphere or contralateral to pain side. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 16 days)</td>
<td>Pain score reduction (mean: -58%) after active but not sham tDCS at the end of stimulation protocol. No significant pain relief at 16 days. No confounding effect on depression or anxiety.</td>
<td>III</td>
</tr>
<tr>
<td>Soler et al., 2010</td>
<td>38 patients with spinal cord injury at various anatomical levels and of various origins (10 active, 9 active + virtual illusion 9 sham, 9 sham + virtual illusion)</td>
<td>Anode: M1 (C3/C4) of the dominant hemisphere or contralateral to pain side. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 10 sessions (2 weeks; FU: 12 weeks)</td>
<td>Greater pain score reduction after active tDCS + virtual illusion, compared to the 3 other groups, up to 12 weeks after stimulation.</td>
<td>III</td>
</tr>
<tr>
<td>Wrigley et al., 2013</td>
<td>10 patients with complete spinal cord injury at thoracic level but of various origins (crossover)</td>
<td>Anode: M1 (C3/C4) of the dominant hemisphere. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 6 months)</td>
<td>No pain relief after active tDCS.</td>
<td>II</td>
</tr>
<tr>
<td>Souto et al., 2014</td>
<td>20 patients with lower limb pain secondary to HTLV-I infection (10 active, 10 sham)</td>
<td>Anode: left M1 (C3); Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; no FU)</td>
<td>Similar global pain relief after active and sham tDCS, but more responders &gt;50% pain relief after active vs. sham tDCS (8/10 vs. 3/10).</td>
<td>II</td>
</tr>
<tr>
<td>Yoon et al., 2014</td>
<td>16 patients with traumatic spinal cord injury at various anatomical levels (10 active, 6 sham)</td>
<td>Anode: left M1 (C3); Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 10 sessions (1 week; FU: 1 day)</td>
<td>Pain score reduction (mean: -23%) after active tDCS, correlated with metabolic changes in the cerebellum, medulla, anterior cingulate and prefrontal cortices.</td>
<td>III</td>
</tr>
<tr>
<td><strong>Recommendation: anodal tDCS of left M1 (or contralateral to pain side) is possibly effective in lower limb pain due to spinal cord lesion (Level C)</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Chronic neuropathic pain of the lower limbs due to peripheral nervous system lesion</strong></td>
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</tr>
<tr>
<td>Kim et al., 2013</td>
<td>40 patients with diabetic polyneuropathy (20 active, 20 sham)</td>
<td>Anode: left M1 (C3); Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 4 weeks)</td>
<td>Greater pain score reduction (mean -34% vs. -14%) after active vs. sham tDCS, up to 4 weeks after stimulation.</td>
<td>II</td>
</tr>
<tr>
<td>Attal et al., 2016</td>
<td>35 patients with unilateral lumbar radiculopathy (23 active, 12 sham)</td>
<td>Anode: left M1 (C3/C4) contralateral to pain side. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 3 sessions (1 week; FU: 5 days)</td>
<td>No difference between active and sham tDCS</td>
<td>II</td>
</tr>
<tr>
<td><strong>No recommendation for anodal tDCS of left M1 (or contralateral to pain side) in lower limb pain due to peripheral nervous system lesion</strong></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Chronic pain related to temporomandibular disorder</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Donnell et al., 2015</td>
<td>24 patients (12 active, 12 sham)</td>
<td>Multipolar tDCS. Two anodes: left M1 (C3+C5). Two cathodes: placed anteriorly (FC3+FC5)</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 1 month)</td>
<td>Increase in the size of pain-free mouth opening one week after tDCS and in the number of responders (&gt;50% pain relief) four weeks after tDCS</td>
<td>III</td>
</tr>
<tr>
<td>Oliveira et al., 2015</td>
<td>32 patients (16 active, 16 sham; combined with cervicomandibular exercises)</td>
<td>Anode: M1 (C3/C4) contralateral to pain side. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 3 weeks)</td>
<td>No difference regarding effect on pain intensity or pressure pain threshold between active and sham tDCS</td>
<td>II</td>
</tr>
<tr>
<td><strong>No recommendation for anodal tDCS of left M1 (or contralateral to pain side) in pain related to temporomandibular disorder</strong></td>
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<tr>
<td><strong>Fibromyalgia</strong></td>
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<tr>
<td>Fregni et al., 2006e</td>
<td>32 patients (11 active M1, 10 sham)</td>
<td>Anode: left M1 (C3); Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 21 days)</td>
<td>Greater pain score reduction (mean: -58% vs. -33%) and quality of life improvement after active vs. sham tDCS, remaining significant up to 21 days.</td>
<td>II</td>
</tr>
<tr>
<td>Valle et al., 2009</td>
<td>41 patients (14 active M1, 14 sham)</td>
<td>Anode: left M1 (C3); Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 10 sessions (2 weeks; FU: 60 days)</td>
<td>Greater pain score reduction (mean: -30% vs. -10%) and quality of life improvement after active vs. sham tDCS, remaining significant up to 60 days.</td>
<td>II</td>
</tr>
</tbody>
</table>
Riberto et al., 2011  
23 patients (11 active, 12 sham; combined with a rehabilitation program)  
Anode: left M1 (C3). Cathode: right supraorbital region  
1 mA, 20 min, 10 sessions (10 weeks; no FU)  
More reduced impact of pain on the quality of life after active vs. sham tDCS, but no differential effect on pain intensity, depression, and anxiety.  
II

Fagerlund et al., 2015  
48 patients (24 active, 24 sham)  
Anode: left M1 (C3). Cathode: right supraorbital region  
2 mA, 20 min, 5 sessions (1 week; FU: 30 days)  
Greater pain score reduction (mean: -14% vs. -2%) and less distress 30 days after active vs. sham tDCS.  
II

Foerster et al., 2015  
12 patients (crossover)  
Anode: left M1 (C3). Cathode: right supraorbital region  
2 mA, 20 min, 5 sessions (1 week; FU: 1 week)  
Pain score reduction (mean: -35%) after active but not sham tDCS, correlated to glutamatergic transmission in the anterior cingulate.  
II

Jales Junior et al., 2015  
20 patients (10 active, 10 sham)  
Anode: left M1 (C3). Cathode: right supraorbital region  
1 mA, 20 min, 10 sessions (10 weeks; 1 session/week; no FU)  
Pain score reduction (mean: -40%) and quality of life improvement after active but not sham tDCS.  
II

Cummiford et al., 2016  
13 patients (crossover)  
Anode: left M1 (C3). Cathode: right supraorbital region  
2 mA, 20 min, 5 sessions (1 week, no FU)  
No difference in pain score reduction between active and sham tDCS but differences in functional cerebral connectivity changes assessed by fMRI.  
III

Recommendation: Anodal tDCS of left M1 is probably effective to relieve pain in fibromyalgia patients (Level B)

FU: follow-up; HTLV-1: Human T-lymphotropic virus type 1; M1: primary motor cortex

Table 2. Repeated tDCS session protocols in motor symptoms of Parkinson’s disease

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benninger et al., 2010</td>
<td>25 patients (HY 2-4) (13 active, 12 sham)</td>
<td>Anode: both (pre)motor and prefrontal cortices. Cathodes: mastoids</td>
<td>2 mA, 20 min, 8 sessions (2.5 weeks; FU: 3 months)</td>
<td>Improvement of gait in off-condition for a short time and of bradykinesia in on- and off-conditions, persisting at least 3 months after the stimulation period.</td>
<td>II</td>
</tr>
<tr>
<td>Valentino et al., 2014</td>
<td>10 patients (HY 2.5-4) (crossover)</td>
<td>Anode: M1 (leg area) contralateral to dominant lower limb. Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 1 month)</td>
<td>Improvement of motor performance (motor UPDRS) and gait, with reduction in number and duration of freezing episodes, persisting at least 4 weeks after the stimulation period.</td>
<td>III</td>
</tr>
<tr>
<td>Costa-Ribeiro et al., 2016a,b</td>
<td>22 patients (HY 1-3) (11 active, 11 sham), combined with cueing gait training</td>
<td>Anode: 2 cm anterior to Cz. Cathode: supraorbital region opposite to the most affected side</td>
<td>2 mA, 20 min, 10 sessions (3 weeks; FU: 1 month)</td>
<td>Improvement of motor performance and gait due to gait training was similar between the active and sham tDCS groups, but the clinical gain and a decrease in motor threshold lasted one month only in the active group.</td>
<td>II</td>
</tr>
</tbody>
</table>

No recommendation for anodal tDCS of the motor cortex in motor symptoms of Parkinson’s disease

FU: follow-up; HY: Hoehn and Yahr disease stage; M1: primary motor cortex; UPDRS: Unified Parkinson’s Disease Rating Scale

Table 3. Repeated tDCS session protocols in motor stroke

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rossi et al., 2013</td>
<td>50 patients (2 days after stroke) (25 active, 25 sham)</td>
<td>Anode: ipsilesional M1 (C3/C4). Cathode: opposite supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 3 months)</td>
<td>No difference in motor or daily living changes (FMA, NIHSS, mRS, BI) after active vs. sham tDCS at the end of the intervention and 3 months later.</td>
<td>I</td>
</tr>
</tbody>
</table>

Anodal stimulation of the ipsilesional motor cortex

Acute phase
Sattler et al., 2015 20 patients (1-14 days after stroke) (10 active, 10 sham; combined with 5Hz radial nerve stimulation) Anode: ipsilesional M1 (TMS hotspot); Cathode: opposite supraorbital region 1.2 mA, 13 min, 5 sessions (1 week; FU: 1 month) Improvement of upper limb motor performance (JHFT, but not FMA) after anodal vs. sham tDCS at 2 and 4 weeks after intervention. Postacute phase

Hesse et al., 2011 56 patients (3-8 weeks after stroke) (28 active, 28 sham; combined with robot-assisted arm motor training) Anode: ipsilesional M1 (C3/C4); Cathode: opposite supraorbital region 2 mA, 20 min, 30 sessions (6 weeks; FU: 3 months) No difference in motor or daily living changes (FMA, BBT, MRC, mAS, BI) after active vs. sham tDCS at the end of the intervention and 3 months later. Postacute phase

Khedr et al., 2013 27 patients (1-4 weeks after stroke) (14 active, 13 sham; combined with conventional physical therapy) Anode: ipsilesional M1; Cathode: opposite supraorbital region 2 mA, 25 min, 6 sessions (1 week; FU: 3 months) Improvement of four limb motor performance and daily living activities (OMCASS, BI), correlated with MT reduction after active vs. sham tDCS at the end of the intervention and 3 months later. Postacute phase

Chang et al., 2015 24 patients (1-4 weeks after stroke) (12 active, 12 sham; combined with robot therapy for upper limb) Anode: ipsilesional M1 (leg area, TMS hotspot); Cathode: opposite supraorbital region 2 mA, 10 min, 10 sessions (2 weeks; FU: 1 day) Improvement of lower limb motor performance (FMA), MEP facilitation, but no effect on gait after anodal vs. sham tDCS. Chronic phase

Geron et al., 2011 30 patients (14-34 months after stroke) (10 active, 10 sham; combined with robot-assisted gait training) Anode: ipsilesional M1 (leg area); Cathode: opposite supraorbital region 1.5 mA, 7 min, 10 sessions (2 weeks; FU: 2 weeks) No difference in walking tests after active vs. sham tDCS. Chronic phase

Viana et al., 2014 20 patients (14-52 months after stroke) (10 active, 10 sham; combined with virtual reality therapy) Anode: ipsilesional M1 (C3/C4); Cathode: opposite supraorbital region 2 mA, 13 min, 15 sessions (5 weeks; no FU) Improvement of quality of life (SSQOL) but no difference in motor performance (FMA, WFMT, mAS) after active vs. sham tDCS Chronic phase

Allman et al., 2016 24 patients (6-141 months after stroke) (11 active, 13 sham; combined with motor training) Anode: ipsilesional M1 (C3); Cathode: opposite supraorbital region 1 mA, 20 min, 9 sessions (2 weeks;FU: 3 months) Improvement of motor performance on ARAT and WFMT but not on upper extremity FMA after active vs. sham tDCS at the end of the intervention and 3 months later Chronic phase

No recommendation for anodal tDCS of the ipsilesional motor cortex in motor stroke

Cathodal stimulation of the contralesional motor cortex

Postacute phase

Hesse et al., 2011 57 patients (3-8 weeks after stroke) (29 active, 28 sham; combined with robot-assisted arm motor training) Anode: opposite supraorbital region; Cathode: contralesional M1 (C3/C4) 2 mA, 20 min, 30 sessions (6 weeks; FU: 3 months) No difference in motor or daily living changes (FMA, BBT, MRC, mAS, BI) after active vs. sham tDCS at the end of the intervention and 3 months later. Postacute phase

Khedr et al., 2013 26 patients (1-4 weeks after stroke) (13 active, 13 sham; combined with conventional physical therapy) Anode: opposite supraorbital region; Cathode: contralesional M1 2 mA, 25 min, 6 sessions (1 week; FU: 3 months) Improvement of four limb motor performance and daily living activities (OMCASS, BI), correlated with MT reduction after active vs. sham tDCS at the end of the intervention and 3 months later. Postacute phase

No recommendation for cathodal tDCS of the contralesional motor cortex in motor stroke

ARAT, action research arm test; BBT: box and block test; BI: Barthel index; FMA: Fugl-Meyer assessment; FU: follow-up; JHFT: Jebsen hand function test; mAS: modified Ashworth scale; MEP: motor evoked potential; MRC: Medical Research Council score; mRS: modified Rankin scale; NIHSS: National Institute of Health stroke score; OMCASS: Orgogozo's MCA scale; SSQOL: stroke specific quality of life scale; TMS: transcranial magnetic stimulation; WFMT: Wolf Motor Function Test

Table 4. Repeated tDCS session protocols in poststroke nonfluent aphasics

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anodal stimulation of Broca’s area</td>
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</tbody>
</table>
### Table 5. Repeated tDCS session protocols in multiple sclerosis fatigue

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation number, intensity, session duration, total number of sessions (protocol follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ferrucci et al., 2014</td>
<td>23 patients (19 RR, 4 SP; EDSS 0-6.5) (crossover)</td>
<td>Anode: bilateral motor cortex / Broca’s area (F5). Cathode: right supraorbital region</td>
<td>1.5 mA, 10 min, 5 sessions (1 week; FU: 3 months)</td>
<td>Improvement vs. sham tDCS at 2 months after the intervention, without correlation with MEP and SEP changes</td>
<td>II</td>
</tr>
<tr>
<td>Tecchio et al., 2014</td>
<td>10 patients (7 RR, 1 SP, 2 PP; EDSS 0-3.5) (crossover)</td>
<td>Anode: whole body S1 personalized electrode. Cathode: right supraorbital region</td>
<td>1.5 mA, 10 min, 5 sessions (1 week; FU: 8 weeks)</td>
<td>No difference in naming accuracy and time after active vs. sham tDCS at the end of the intervention and 3 months later, but higher effect sizes in naming time after active tDCS.</td>
<td>II</td>
</tr>
<tr>
<td>Tecchio et al., 2015</td>
<td>13 patients (13 RR; EDSS 0-3.5) (crossover)</td>
<td>Anode: whole body S1 personalized electrode. Cathode: right supraorbital region</td>
<td>1.5 mA, 10 min, 5 sessions (1 week; FU: no FU)</td>
<td>Improvement in picture description, noun and verb naming after active tDCS at the end of the intervention, with difference according to the integrity of different left subcortical structures</td>
<td>II</td>
</tr>
</tbody>
</table>

No recommendation for anodal tDCS of sensory or motor cortex in multiple sclerosis fatigue

EDSS: Expanded Disability Status Scale; FU: follow-up; MEP: motor evoked potentials; mFIS: modified fatigue impact scale; PP: primary progressive; RR: relapsing-remitting; S1: primary somatosensory cortex; SEP: somatosensory evoked potentials; SP: secondary progressive

### Table 6. Repeated tDCS session protocols in Alzheimer’s disease

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cotelli et al., 2014a</td>
<td>36 patients (mean MMSE 20-22) (12 active + memory training, 12 active + motor training, 12 sham + memory training)</td>
<td>Anode: left DLPFC. Cathode: right deltoid muscle</td>
<td>2 mA; 25 min, 10 sessions (2 weeks; FU: 6 months)</td>
<td>No difference in cognitive performance improvement (face-name association task) after active vs. sham tDCS combined with memory training, but no improvement when combined with motor training.</td>
<td>II</td>
</tr>
<tr>
<td>Khedr et al., 2014b</td>
<td>34 patients (MMSE 12-23) (11 anodal, 12 cathodal, 11 sham)</td>
<td>Anode: left DLPFC. Cathode: right supraorbital region</td>
<td>2 mA; 25 min, 10 sessions (2 weeks; FU: 2 months)</td>
<td>Improvement in cognitive performance (MMSE) after either anodal or cathodal vs. sham tDCS at 2 months after the intervention</td>
<td>II</td>
</tr>
</tbody>
</table>
(or the reverse) intervention, with reduction of P300 latency, but no effect on M1 excitability.

Suemoto et al., 2014
40 patients (MMSE 10-20) (20 active, 20 sham) Anode: left DLPFC, Cathode: right supraorbital region
2 mA, 20 min, 6 sessions (2 weeks; FU: 1 week) No change in apathy scores, global cognition, and neuropsychiatric symptoms after active vs. sham tDCS. II

Bystad et al., 2016
25 patients (MMSE >17) (12 active, 13 sham) Anode: left temporal lobe (T3). Cathode: right supraorbital region
2 mA, 30 min, 6 sessions (2 weeks; no FU) No change in verbal memory function after active vs. sham tDCS. II

No recommendation for anodal tDCS of the left DLPFC in Alzheimer’s disease

DLPFC: dorsolateral prefrontal cortex; FU: follow-up; M1: primary motor cortex; MMSE: Mini Mental State Examination

Table 7. Repeated tDCS session protocols in tinnitus

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shekhawat et al., 2013a</td>
<td>40 patients (20 active, 20 sham; followed by sound therapy from hearing aid use for 6 months) Anode: left temporoparietal cortex (between C3 and T5). Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 6 months)</td>
<td>No difference in the effect on tinnitus functional index, handicap questionnaire, and severity scores after active vs. sham tDCS.</td>
<td>II</td>
<td></td>
</tr>
<tr>
<td>Teismann et al., 2014</td>
<td>32 patients (10 anodal, 11 cathodal, 11 sham; combined with tailor-made notched music training) Anode: left temporoparietal cortex / Heschl’s Gyrus (between C3 and T3, 1 cm inferior). Cathode: right supraorbital region</td>
<td>2 mA, 30 min, 5 sessions (1 week; FU: 31 days)</td>
<td>No difference in the effect on tinnitus handicap questionnaire and inventory after either anodal or cathodal vs. sham tDCS.</td>
<td>II</td>
<td></td>
</tr>
<tr>
<td>Forogh et al., 2016</td>
<td>22 patients (11 active, 11 sham) Anode: left temporoparietal cortex (between C3 and T5). Cathode: right supraorbital region</td>
<td>2 mA, 20 min, 5 sessions (1 week; FU: 2 weeks)</td>
<td>No difference in the effect on tinnitus handicap inventory and VAS after either active vs. sham tDCS.</td>
<td>II</td>
<td></td>
</tr>
<tr>
<td>Hyvärinen et al., 2016</td>
<td>19 patients (10 left temporal, 9 sham) Anode: left temporal cortex. Cathode: right frontal region</td>
<td>2 mA, 20 min, 10 sessions (10 consecutive days; FU: 4 weeks after treatment start)</td>
<td>No difference in the effect on tinnitus handicap inventory after active vs. sham tDCS.</td>
<td>II</td>
<td></td>
</tr>
</tbody>
</table>

Recommendation: anodal tDCS of the left temporoparietal cortex is probably ineffective in chronic tinnitus (Level B)

FU: follow-up; VAS: visual analogue scale

Table 8. Repeated tDCS session protocols in depression

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anodal tDCS of the left DLPFC with right orbitofrontal cathode</td>
<td></td>
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</tr>
<tr>
<td>Boggio et al., 2008a</td>
<td>40 patients (non-medicated MDD) (21 active DLPFC, 9 active occipital, 10 sham) Anode: left DLPFC (F3) or occipital cortex (Oz). Cathode: right orbitofrontal region</td>
<td>2 mA, 20 min, 10 sessions (2 weeks; FU: 30 days)</td>
<td>Mood improvement (HDRS, BDI) after active vs. sham tDCS of the left DLPFC.</td>
<td>II</td>
<td></td>
</tr>
<tr>
<td>Loo et al., 2010</td>
<td>34 patients (MDD, including 20 non-medicated, MADRS &gt;20) (19 active, 15 sham) Anode: left DLPFC (F3). Cathode: right orbitofrontal region</td>
<td>1 mA, 20 min, 5 active or sham sessions (1.5 week), followed by 5 active sessions (1.5 week; FU: 1 month)</td>
<td>No difference in mood improvement (HDRS-17, MADRS) after active vs. sham tDCS.</td>
<td>II</td>
<td></td>
</tr>
<tr>
<td>Loo et al., 2012</td>
<td>60 patients (MDD, including 17 non-medicated and 8 bipolars, MADRS &gt;20) (31 active, Anode: left DLPFC (F3). Cathode: right orbitofrontal region (F8)</td>
<td>2 mA, 20 min, 15 active or sham sessions (3 weeks), followed by 15 active</td>
<td>Mood improvement (MADRS) after active vs. sham tDCS, but no difference in responder rate (13%).</td>
<td>I</td>
<td></td>
</tr>
</tbody>
</table>
Mondino et al., 2016
Fröhlich et al., 2015
Mondino et al., 2014
Brunelin et al., 2012a
Allden et al., 2012
Blumberger et al., 2015
Bennabi et al., 2013b
Brunoni et al., 2012
Bennabi et al., 2012
Palm et al., 2013

### Anodal tDCS of the left DLPFC with right DLPFC cathode

**Recommendation:** anodal tDCS of the left DLPFC with right orbitofrontal cathode is probably effective in patients with no drug-resistant major depressive episode (Level B) and probably ineffective in patients with drug-resistant major depressive episode (Level B).

**Anodal tDCS of the left DLPFC with right DLPFC cathode**

**Table 9. Repeated tDCS session protocols in schizophrenia**

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Stimulation intensity, session duration, total number of sessions (protocol duration; follow-up)</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brunelin et al., 2012a</td>
<td>30 patients with AVH (15 active, 15 sham)</td>
<td>Anode: left DLPFC (between F3 and FP1). Cathode: left temporoparietal junction (between T3 and P3)</td>
<td>2 mA, 20 min, 10 sessions (1 week; FU: 3 months)</td>
<td>Greater AVH reduction (mean (AHRS): -31% vs. -8%) after active vs. sham tDCS, remaining significant up to 3 months. Improvement of other schizophrenia symptoms, including negative symptoms (PANSS) after active vs. sham tDCS at the end of the intervention.</td>
<td>II</td>
</tr>
<tr>
<td>Fitzgerald et al., 2014</td>
<td>24 patients (11 bilateral, 13 unilateral)</td>
<td>Anode: left DLPFC (F3). Cathode: left temporoparietal junction (TP3) (unilaterally or bilaterally F3+FP3). Return reference: C2</td>
<td>2 mA, 20 min, 15 sessions (3 weeks; no FU)</td>
<td>No difference in any schizophrenia symptoms (including AVH and negative symptoms) after active vs. sham tDCS at the end of the intervention.</td>
<td>III</td>
</tr>
<tr>
<td>Mondino et al., 2015</td>
<td>28 patients with AVH (15 active, 13 sham)</td>
<td>Anode: left DLPFC (between F3 and FP1). Cathode: left temporoparietal junction (between T3 and P3). Return reference: C2</td>
<td>2 mA, 20 min, 5 sessions (1 week; no FU)</td>
<td>Greater AVH frequency reduction (mean: -46% vs. +7.5%) after active vs. sham tDCS. Reduction in AVH frequency correlated with improvement in source monitoring performances (decrease of externalization bias).</td>
<td>II</td>
</tr>
<tr>
<td>Frohlich et al., 2016</td>
<td>26 patients with AVH (13 active, 13 sham)</td>
<td>3-electrode montage. Anode: left DLPFC (between F3 and FP1). Cathode: left temporoparietal junction (between T3 and P3). Return reference: Cz</td>
<td>2 mA, 20 min, 5 sessions (1 week; no FU)</td>
<td>No difference in AVH reduction, (mean (AHRS): -24% vs. -34%) after active vs. sham tDCS. No effect on other symptoms.</td>
<td>II</td>
</tr>
<tr>
<td>Mondino et al., 2016</td>
<td>23 patients with AVH (11 active, 12 sham)</td>
<td>Anode: left DLPFC (between F3 and FP1). Cathode: left temporoparietal junction (between T3 and P3)</td>
<td>2 mA, 20 min, 10 sessions (1 week; no FU)</td>
<td>Greater AVH reduction (mean (AHRS): -28% vs. -10%) after active vs. sham tDCS, correlated with a reduction of functional connectivity between the left</td>
<td>II</td>
</tr>
</tbody>
</table>
temporo-parietal junction and anterior insula.

No recommendation for anodal tDCS of the left DLPFC with left temporo-parietal cathode in schizophrenia

Anodal tDCS of the left DLPFC with right supraorbital cathode

Smith et al., 2015
29 patients who were cigarette smokers (14 active, 15 sham)
Anode: left DLPFC (F3).
Cathode: right supraorbital (Fp2-A8)
2 mA, 20 min, 5 sessions (1 week; no FU)
Improvement in various scores of the MATRICS Consensus Cognitive Battery after active vs. sham tDCS, but no difference in PANSS scores, AVH, and cigarette craving between the two conditions.
II

Palm et al., 2016b
20 patients with negative symptoms (10 active, 10 sham)
Anode: left DLPFC (F3).
Cathode: right supraorbital (Fp2-A8)
2 mA, 20 min, 10 sessions (2 weeks; FU: 2 weeks)
Reduction of SANS and PANSS total scores after active vs. sham tDCS.
II

No recommendation for anodal tDCS of the left DLPFC with right supraorbital cathode in schizophrenia

AHRS: auditory hallucinations rating scale; AVH: auditory verbal hallucinations; DLPFC: dorsolateral prefrontal cortex; FU: follow-up; PANSS: positive and negative symptoms scale; SANS: Scale for the Assessment of Negative Symptoms

Table 10. Repeated tDCS session protocols in addiction/craving

<table>
<thead>
<tr>
<th>Articles</th>
<th>Number and type of patients (protocol design)</th>
<th>Stimulation electrode location</th>
<th>Clinical results</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol</td>
<td></td>
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<tr>
<td>Klauss et al., 2014</td>
<td>33 alcoholics (16 active, 17 sham)</td>
<td>Anode: right DLPFC (F4). Cathode: left DLPFC (F3)</td>
<td>2 mA, 13 min, 10 sessions (1 week; FU: 6 months)</td>
<td>II</td>
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<tr>
<td></td>
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<td></td>
<td>Improvement in overall perception of quality of life after active vs. sham tDCS, but no difference in craving scores, frontal function, global mental status, and anxiety-depression symptoms. At 6 months, alcohol-abstinent subjects were more numerous after active vs. sham tDCS (8/16 vs/ 2/17).</td>
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<tr>
<td>Crack-Cocaine</td>
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<tr>
<td>Batista et al., 2015</td>
<td>36 crack-cocaine users (17 active, 19 sham)</td>
<td>Anode: right DLPFC (F4). Cathode: left DLPFC (F3)</td>
<td>2 mA, 20 min, 5 sessions (6 weeks; FU: 1 week)</td>
<td>II</td>
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<td></td>
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<td></td>
<td>Improvement in craving scores, anxiety, and overall perception of quality of life after active vs. sham tDCS. In addition, craving scores decreased linearly over 4 weeks.</td>
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<tr>
<td>Smoking</td>
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<tr>
<td>Boggio et al., 2009b</td>
<td>27 smokers (13 active, 14 sham)</td>
<td>Anode: right DLPFC (F4). Cathode: left DLPFC (F3)</td>
<td>2 mA, 20 min, 5 sessions (1 week; no FU)</td>
<td>II</td>
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<tr>
<td></td>
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<td></td>
<td>Decrease in the number of cigarettes smoked after active vs. sham tDCS in the week of intervention. In addition, cumulative reduction of craving for cue- provoked smoking after active tDCS</td>
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</tr>
<tr>
<td>Fecteau et al., 2014</td>
<td>12 smokers (crossover)</td>
<td>Anode: right DLPFC (F4). Cathode: left DLPFC (F3)</td>
<td>2 mA, 30 min, 5 sessions (1 week; FU: 4 days)</td>
<td>II</td>
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<tr>
<td></td>
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<td></td>
<td>Decrease in the number of cigarettes smoked after active vs. sham tDCS, lasting up to 4 days after the end of the intervention. Smokers rejected more offers of cigarettes with reward sensitive effects after active tDCS</td>
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<tr>
<td>Recommendation: combined anodal tDCS of the right DLPFC and cathodal tDCS of the left DLPFC is probably effective in addiction/craving (Level B)</td>
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</tbody>
</table>

Table 11. Summary of recommendations on tDCS efficacy according to clinical indication

<table>
<thead>
<tr>
<th>Clinical indication</th>
<th>tDCS protocol</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic neuropathic pain of the lower limbs due to spinal cord lesion</td>
<td>Anodal tDCS of left M1 (or contralateral to pain side) is possibly effective in case of spinal cord lesion (Level C). No recommendation in case of peripheral nervous system lesion</td>
<td></td>
</tr>
<tr>
<td>Chronic non-neuropathic oral, facial, low back, abdominal, or pelvic pain</td>
<td>No recommendation for anodal tDCS of left M1 (or contralateral to pain side)</td>
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<tr>
<td>Fibromyalgia</td>
<td>Anodal tDCS of left M1 is probably effective (Level B)</td>
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<tr>
<td>Migraine</td>
<td>No recommendation for anodal tDCS of the left M1 or cathodal tDCS of V1</td>
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<tr>
<td>Postoperative pain</td>
<td>No recommendation for anodal tDCS of M1 or the left DLPFC</td>
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<tr>
<td>Parkinson’s disease (motor symptoms)</td>
<td>No recommendation for anodal tDCS of the motor cortex</td>
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<tr>
<td>Condition (cognitive symptoms)</td>
<td>No recommendation for anodal tDCS of the left DLPFC</td>
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<tr>
<td>Dystonia (focal or generalized)</td>
<td>No recommendation for anodal or cathodal tDCS of the motor cortex</td>
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<tr>
<td>Motor stroke</td>
<td>No recommendation for anodal tDCS of the ipsilesional motor cortex in acute, postacute, or chronic stage</td>
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<tr>
<td></td>
<td>No recommendation for cathodal tDCS of the contralesional motor cortex in postacute stage</td>
<td></td>
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<tr>
<td></td>
<td>No recommendation for bihemispheric tDCS of the motor cortex (ipsilesional anode + contralesional cathode) in chronic stage</td>
<td></td>
</tr>
<tr>
<td>Nonfluent poststroke aphasia</td>
<td>No recommendation for anodal tDCS of left Broca’s area or Wernicke’s area or cathodal tDCS of their right homologous region in postacute or chronic stage</td>
<td></td>
</tr>
<tr>
<td>Multiple sclerosis (fatigue or cognitive disorders)</td>
<td>No recommendation for anodal tDCS of sensory or motor cortex or the left DLPFC</td>
<td></td>
</tr>
<tr>
<td>Multiple sclerosis (sensory or motor symptoms)</td>
<td>No recommendation for anodal tDCS of M1</td>
<td></td>
</tr>
<tr>
<td>Epilepsy</td>
<td>No recommendation for cathodal tDCS of the epileptic focus or anodal tDCS of the left DLPFC</td>
<td></td>
</tr>
<tr>
<td>Disorders of consciousness</td>
<td>No recommendation for anodal tDCS of the left DLPFC</td>
<td></td>
</tr>
<tr>
<td>Alzheimer’s disease</td>
<td>No recommendation for anodal tDCS of the left DLPFC or the temporoparietal cortex</td>
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<tr>
<td>Tinnitus</td>
<td>Anodal tDCS of the left temporoparietal cortex is probably ineffective (Level B)</td>
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<tr>
<td></td>
<td>No recommendation for bihemispheric tDCS of the DLPFC (right anode + left cathode)</td>
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<tr>
<td>Depression</td>
<td>Anodal tDCS of the left DLPFC with right orbitofrontal cathode is probably effective in non-drug-resistant major depression (Level B) and probably ineffective in drug-resistant major depression (Level B)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No recommendation for bihemispheric tDCS of the DLPFC (left anode + right cathode)</td>
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<tr>
<td>Schizophrenia (AVH or negative symptoms)</td>
<td>No recommendation for anodal tDCS of the left DLPFC with left temporoparietal or right supraorbital cathode</td>
<td></td>
</tr>
<tr>
<td>Addiction / craving</td>
<td>Bihemispheric tDCS of the DLPFC (right anode + left cathode) is probably effective in addiction/craving (alcohol, drugs, smoking) (Level B)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No recommendation for anodal tDCS of the left DLPFC with right supraorbital cathode</td>
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</tbody>
</table>

“No recommendation” means the absence of sufficient evidence to date, but not the evidence for an absence of effect.