Non-invasive brain stimulation for improving gait, balance, and lower limbs motor function in stroke

Jitka Veldema1,2* and Alireza Gharabaghi2

Abstract
Objectives: This systematic review and meta-analysis aim to summarize and analyze the available evidence of non-invasive brain stimulation/spinal cord stimulation on gait, balance and/or lower limb motor recovery in stroke patients.

Methods: The PubMed database was searched from its inception through to 31/03/2021 for randomized controlled trials investigating repetitive transcranial magnetic stimulation or transcranial/trans-spinal direct current/alternating current stimulation for improving gait, balance and/or lower limb motor function in stroke patients.

Results: Overall, 25 appropriate studies (including 657 stroke subjects) were found. The data indicates that non-invasive brain stimulation/spinal cord stimulation is effective in supporting recovery. However, the effects are inhomogeneous across studies: (1) transcranial/trans-spinal direct current/alternating current stimulation induce greater effects than repetitive transcranial magnetic stimulation, and (2) bilateral application of non-invasive brain stimulation is superior to unilateral stimulation.

Conclusions: The current evidence encourages further research and suggests that more individualized approaches are necessary for increasing effect sizes in stroke patients.

Keywords: rTMS, tDCS, tACS, tsDCS, Stroke, Gait, Balance, Lower limb function

Introduction
Each year, approximately 795,000 people experience a new or recurrent stroke [1]. Walking and balance disturbances are common post-stroke complications, affecting about two-thirds of stroke survivors [2]. These deficits are associated with worsened quality of life, impeded community reintegration [3], and an increased risk of falling [4]. The ability to walk independently is the most common rehabilitation goal after stroke [5]. However, about 50% of stroke survivors suffer from an impaired walking ability 6 months after current standard of care [2]. Other interventions are therefore needed to improve recovery. Thus, the development of innovative therapeutic strategies for improving balance and walking ability is one of the top research priorities in stroke rehabilitation [6]. Non-invasive neuromodulation methods such as repetitive transcranial magnetic stimulation (rTMS), transcranial direct/alternating current stimulation (tDCS/tACS) and trans-spinal direct current stimulation (tsDCS) can modulate neural processing and have thus the potential to counteract maladaptive neural plasticity after stroke and contribute to a better recovery [7, 8].

Neural background of walking and balance
Neuroimaging studies have shown that walking and balance are complex sensorimotor functions controlled by integrated cortical, subcortical, and spinal networks...
A single photon emission computed tomography study demonstrates bilateral activation within the primary sensorimotor area, supplementary motor area, basal ganglia as well as within the visual cortex, cerebellar vermis, and part of the left lower temporal lobe during walking [12]. Similarly, a positron emission tomography study shows a bilateral increase of cerebral blood flow within the primary sensory cortex, primary motor cortex and supplementary motor cortex as well as within the anterior part of the cerebellum during active and passive bipedal movement [13]. A recent meta-analysis indicates a key role of the brainstem, cerebellum, basal ganglia, thalamus, and several cortical regions during postural control [9]. Accordingly, another meta-analysis shows that the cerebellum, basal ganglia, thalamus, hippocampus, inferior parietal cortex, and frontal lobe regions are involved during balance tasks [14]. Importantly, the available data indicates that also a spinal network may be involved in postural balance and gait control [15]. E.g., multiple studies demonstrate that balance training induces suppression of H-reflexes [16]. Thus, it is conceivable that the application of non-invasive brain stimulation over several cortical regions as well as over the cerebellum, the brainstem and the spinal cord may be effective in the modulation of walking, balance and/or lower limbs motor function.

**Stroke-induced changes of neural control during walking**

Up to now, different neuroimaging techniques have been used to investigate the neural mechanism of walking disability and walking recovery in stroke patients. A large part of the available data demonstrates a stroke-induced disinhibition of the contralesional hemisphere with a shift of the between-hemispheric balance to the detriment of the affected hemisphere, as well as a correlation between normalization of neural processing and favorable motor recovery [17–21]. A diffusion tensor MRI demonstrates a between-hemispheric asymmetry in fractional anisotropy of the posterior limb of the internal capsule [18]. The shift of balance towards the non-lesioned hemisphere correlates with the amount of walking disability [18]. An optical imaging study shows a between-hemispheric imbalance of oxygenated hemoglobin level in the medial primary sensorimotor cortex that is greater in the unaffected hemisphere than in the affected hemisphere. A reduction of this asymmetry is associated with a favorable gait recovery [19]. A TMS study shows an interhemispheric asymmetry of corticomotor excitability of the legs to the detriment of the affected hemisphere, as well as a correlation between the reduction of this asymmetry and a favorable motor outcome [20]. Another TMS trial reveals increased connectivity between the contralesional hemisphere and the affected lower limb, which correlated with the amount of walking disability [18]. A diffusion-weighted MRI shows that the higher the anatomical connectivity between the ipsilesional M1 and the (a) cerebral peduncle, (b) thalamus, and (c) red nucleus, the better is the lower limb motor performance [21]. Furthermore, stroke-related disturbances of the spinal system were detected, as well as its relationship to gait disability. In fact, stroke patients show an increase of the H-reflex, in comparison to healthy subjects [22], and its normalization to be associated with a successful motor recovery of the walking ability [23].

**Non-invasive brain stimulation for network modulation**

The application of non-invasive brain stimulation in rehabilitation aims at prolonged effects on the neural network. It is assumed that these techniques modulate synaptic connectivity, similar to long-term potentiation and long-term depression, which are considered relevant mechanism of plastic reorganization [24]. The amount and the duration of the induced neurophysiological changes depend on the stimulation intensity and duration [25, 26]. The available data indicates that a direct current of at least 0.6 mA that is applied for at least three minutes is sufficient to modulate cortical excitability beyond the stimulation period. Applying tDCS of 1 mA for five to seven minutes leads to short-term changes of cortical excitability that last 10–15 min after the end of stimulation. For long-term modulation of cortical excitability (1 h or more) a current of 1 mA need to be applied over a period of at least 11 min [25]. A single session of rTMS induces cortical excitability changes that last for at least 30 min after the end of stimulation [26]. In previous decades specific certain protocols have been established as either “facilitatory” or “inhibitory” for both rTMS and tDCS/tACS/tsDCS techniques. High-frequency rTMS (≥5 Hz), intermittent theta burst stimulation (iTBS), paired pulse rTMS (inter-stimulus interval 1.5 ms) and anodal tDCS/tACS are considered to have “up-regulating” effects on neural processing. In contrast low-frequency rTMS (1 Hz), continuous theta burst stimulation (cTBS), paired-pulse rTMS (inter-stimulus interval 3 ms) and cathodal tDCS are expected to induce a “down-regulation” [7, 8]. Indeed, several studies demonstrated a modulation of neural processing outside this framework [27–30]. An earlier study has shown that several rTMS protocols (1 Hz, 10 Hz, 15 Hz, 20 Hz) can induce an increase and a decrease of corticospinal excitability in the stimulated hemisphere [27]. Similarly, a more recent trial demonstrated inhibitory and facilitatory influences on corticospinal excitability of both iTBS and cTBS [28]. The TBS-induced effect was highly correlated with the pre-interventional MEP latency [28]. Similarly, cTBS decreased and increased corticospinal
excitability, and its effects correlated with pre-interventional MEP variability and late I-wave recruitment [29]. Also, both anodal and cathodal tDCS induced increases and decreases of corticospinal excitability in the stimulated hemisphere, and the intervention-induced effects correlated with the pre-interventional MEP-latency [30]. Thus, present data shows that the responses to “up” and “down” regulating brain stimulation protocols are inconsistent, already in the healthy condition, and the factors influencing this inter-individual variability are not completely understood.

Another relevant issue is the state-dependency of stimulation effects. Specifically, brain state-dependent single-pulse TMS that was controlled by volitional modulation of sensorimotor beta-band oscillatory activity induced a robust increase of corticospinal excitability [31]. By contrast, the identical stimulation pattern applied independent of the brain state resulted in its decrease. The same was true, when single-pulse TMS was paired with peripheral stimulation; this pairing led to an increase or decrease of corticospinal excitability, when applied during volitional modulation of the sensorimotor beta-band activity or independent of the brain state, respectively [32, 33].

There might be discrepancies, however, between online (i.e., during the intervention) and offline (i.e., after the intervention) stimulation effects, particularly, when the former is applied during behavioral tasks and the latter is done at rest. In this context, a meta-analysis detected timing- and cohort-dependent effects of anodal tDCS on the modulation of working memory. Healthy subjects demonstrated significant offline improvement but no online effects. By contrast, neuropsychiatric patients showed improved working memory during the stimulation but not afterwards [34]. Similarly, another meta-analysis detected timing-dependent effects of rTMS with regard to episodic memory, which was improved or deteriorated, when the stimulation was applied before or during the task [35]. Along the same lines a recent meta-analysis showed that rTMS and tDCS modulated visuospatial abilities in healthy subjects to a larger extent when applied before than during the task [36]. Therefore, we may assume that the effectiveness of non-invasive brain stimulation in supporting motor recovery after stroke will not depend on the stimulation protocol only, but also on the behavioral context.

Non-invasive brain/spinal cord stimulation for improving walking, balance and lower limb motor function in stroke patients
Despite its limitations, the theory of interhemispheric imbalance and rivalry provides the most often used theoretical framework for the application of noninvasive brain stimulation in stroke rehabilitation [37]. Previous studies have demonstrated that several stroke-induced deficiencies, such as upper limb impairment or visuospatial disabilities may be successfully restored by application of non-invasive brain stimulation within this concept [38, 39]. The available data (see previous subchapter) indicates that a similar application may also be useful for supporting gait, balance, and lower limbs’ motor functioning. This means either “inhibitory” stimulation of the contralesional hemisphere or “facilitatory” stimulation of the ipsilesional hemisphere. Furthermore, it is conceivable that “inhibitory” spinal stimulation may be beneficial.

This systematic review and meta-analysis summarize the current evidence for non-invasive brain and spinal cord stimulation to support gait, balance and/or lower limbs function in stroke patients. The effectiveness is analyzed with regard to the technique used (rTMS, tDCS/tACS/tsDCS), protocols applied (anodal/cathodal/bilateral tDCS, low-frequency/high-frequency rTMS, iTBS/cTBS), stimulated hemisphere (affected/non-affected/bilateral), stimulated area (primary motor cortex, cerebellum, supplementary motor area, spinal cord) and applied study design (stimulation amount, evaluation schedule). Furthermore, information regarding the participants (time since stroke, gender, stroke type and location), the exact stimulation location, and methodological quality of the trial is included.

Methods
The protocol of this systematic review and meta-analysis bases on the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) guidelines. A previous registration of the protocol was not performed.

Search protocol
Relevant studies were identified by searching of electronic database PubMed from inception to 31/03/2021. The combination of following search terms was used: (1) “tDCS” or “rTMS” or “tACS” or “tsDCS” and (2) “balance” or “postural control” or “gait” or “walking” or “lower limbs” and (3) stroke. The screening was performed by two independent reviewers (JV and AG). Disagreements were resolved by consensus. Figure 1 illustrates the search strategy based on the PRISMA guidelines.

Eligibility criteria
Trials matching the following criteria were enrolled: (1) randomized controlled trials, (2) diagnosis stroke, (3) at least five participants per intervention, (4) rTMS, tDCS, tACS and/or tsDCS as intervention, (5) pre- and post-interventional assessments of gait, balance and/or lower limb motor function, (6) written in English or German.
Data extraction
The data on gait, balance and/or lower limb function were extracted from the included publications. Depending on the availability, we considered either (1) the pre-intervention, post-intervention and follow up data, or (2) the difference between the pre-intervention and post-intervention data and the difference between the pre-intervention and follow up-data were obtained. The secondary data extracted were: (1) patients’ characteristics (number, gender, age, time since incident, stroke etiology and location), (2) stimulation parameters (technique used, positioning, duration, intensity, number of sessions), (3) methods (study design, assessments, evaluations schedule).

Data synthesis
Based on the methodological approach, the included experiments were split up into (i) experiments comparing real brain stimulation with sham stimulation and (ii) experiments comparing two different brain stimulation protocols. Within this framework, a subcategorization was done depending on (a) stimulation technique (rTMS, tDSC/tACS/tsDCS), (b) stimulation positioning (affected hemisphere, non-affected hemisphere, bilateral, spinal) and (c) stimulation protocol (“facilitatory”: anodal tDSC/tACS/tsDCS, high-frequency rTMS, iTBS, “inhibitory”: cathodal tDSC/tACS/tsDCS, low-frequency rTMS, cTBS, and “combined”).

Statistical analysis
Effect size calculators were used to estimate the effect size and the 95% confidence interval for each experiment [40, 41]. Depending on availability, the calculations are based on either means and standard deviations of pre-intervention, post-intervention and follow up data, or on means and standard deviations of the difference between
the pre-intervention and post-intervention data and the difference between the pre-intervention and follow-up data. Where multiple assessments were applied, the effect sizes and the 95% confidence intervals were calculated for all outcomes, and on their basis average values were determinate for each experiment. The Cohen definition of effect size was used for interpretation (d ≥ 0.2 “small”, d ≥ 0.5 “medium”, d ≥ 0.8 “large”) [42]. The inconsistency test (I²) was applied to evaluate the homogeneity across experiments, where values above 50% indicate high heterogeneity [43].

**Methodological quality assessment**

11-items PEDro scale was applied to evaluate the methodological quality of the studies, such as random allocation, subjects’ and assessors’ blinding, dropout rate etc [44]. The higher the total score, the higher the methodological quality (10–9 excellent, 8–6 good, 5–4 fair and <4 poor).

**Results**

**TDCS, tADCS, tsDCS**

In total, 16 studies tested the effects of tDCS, tACS or tsDCS (Table 1) [45–60]. Their methodological quality varied between fair and excellent (Table 2).

**Participants:** Overall 445 stroke patients (299 males, 146 females) were investigated. The cohorts were inconsistent in terms of the mean time since stroke (between two days and ten years), stroke etiology (309 ischemic, 55 hemorrhagic, 81 na), lesion location (91 subcortical, 109 cortical, 245 na) and lesion site (108 right, 86 left, 251 na).

**Study design:** 14 studies investigated the effectiveness of real tDCS/tACS/tsDCS in comparison to sham stimulation [45–53, 56–60]. Five studies compared different stimulation protocols [45, 54–56, 60]. The overall duration of active stimulation varied between 15 and 400 min (one to 16 sessions with a duration of seven to 40 min were applied). All studies performed a pre- and post-evaluation of the parameters assessed. Nine studies performed additional follow-up evaluations, one up to 24 weeks after completing the intervention [45, 46, 49, 51, 52, 54–56, 58].

**Stimulation protocol:** Seven studies applied anodal tDCS over the ipsilesional hemisphere [45, 47, 49, 52, 58, 60]. Five trials investigated bilateral tDCS (combining anodal tDCS over the ipsilesional hemisphere and cathodal tDCS over the contralesional hemisphere) [45, 50, 57, 59] or bilateral tACS [51]. The contralesional hemisphere alone was stimulated in only two studies – in one study with anodal [60] and in another with cathodal tDCS [45]. One study combined anodal tDCS over supplementary motor area with cathodal tDCS over cerebellum [53]. One trial investigated cathodal tsDCS [56].

**Effectiveness—active stimulation versus sham:** In total, the post-interventional data show a medium-sized effect of tDCS/tACS/tsDCS on gait, balance, and lower limb motor function in stroke patients (Fig. 2). However, effects were inhomogeneous across the studies. Cathodal tDCS over the contralesional hemisphere and bilateral tDCS induce large effects on the evaluated parameters. In contrast, only small effects were found for anodal tDCS/tACS over the ipsilesional hemisphere and for tsDCS. No effect was induced by anodal tDCS over the contralesional hemisphere and by fronto-parietal tDCS.

The follow-up data demonstrate a large-sized effect of stimulation on the assessed parameters, as well as inhomogeneity of the effects across the experiments (Fig. 3). Cathodal tDCS over the contralesional hemisphere and bilateral tDCS evoke large effects. A medium-sized effect was detected for anodal tDCS/tACS over the ipsilesional hemisphere. No effect was found for tsDCS.

**Effectiveness—comparison of different stimulation protocols:** Both the post-interventional and the follow-up data demonstrate that different stimulation protocols may induce significantly different effects on gait, balance, and lower limbs motor function in stroke patients (Figs. 4, 5). In particular bilateral tDCS/tACS and cathodal tDCS over the contralesional hemisphere are superior to other stimulation protocols.

**rTMS**

Nine studies tested rTMS for improving gait, balance, and lower limb motor function in stroke patients (Table 3) [20, 61–68]. Their methodological quality was good to excellent (Table 2).

**Participants:** In total, 212 patients (133 males, 70 females, 9 gender na) were enrolled. The study cohorts were inconsistent regarding the mean time since stroke (11 days to 29 months), stroke etiology (144 ischemic, 44 hemorrhagic, 24 na), stroke location (78 subcortical, 32 cortical, 23 brainstem, 79 na) and lesioned site (95 right, 85 left, 32 na).

**Study design:** All studies investigated the effectiveness of rTMS in comparison to sham stimulation. A direct comparison of different stimulation protocols was not performed. The duration of the total amount of active stimulation varied between 15 and 330 min (between five and 13 sessions with a duration of seven to 30 min were performed). All trials applied pre- and post-interventional assessments. Additional follow-up evaluations, over one week to three months after completing the stimulation, were performed in six trials [61–64, 66, 68].

**Stimulation protocol:** Four studies applied low-frequency rTMS over the contralesional hemisphere [62, 63, 66, 68]. Three trials performed bilateral stimulation (combining high-frequency rTMS or iTBS over the
| References       | Participants number, gender, age/time since stroke | Stroke etiology/ location/affected hemisphere | Stimulation technique/ intensity | Electrodes positioning | Study design/ electrodes positioning technique | Number and duration of sessions/ evaluations schedule | Outcomes                |
|------------------|--------------------------------------------------|-----------------------------------------------|---------------------------------|------------------------|-----------------------------------------------|------------------------------------------------------|-------------------------|
| Bornheim et al. [46] | 31 males, 15 females s/63 ± 12 years/2 ± 0 days | 46 ischemic/na/26 right, 20 left               | (1) 1.0 mA anodal tDCS           | IL C3/C4               | CL Fp2/Fp1                                    | 20 sessions à 20 min/pre, post, two-week, three-month, six-month, and one-year follow-up | FMA-LE                  |
| Chang et al. [48]  | 15 males, 9 females /63 ± 11 years/16 ± 6 days  | 24 ischemic/na/13 right, 11 left               | (1) 2.0 mA anodal tDCS           | IL tibialis anterior M1 | CL SO                                         | 10 sessions à 10 min/pre, post                         | FAC, BBS, MI-LE, FMA-LE, gait (cadence, speed, stride length, step length, step time) |
| Tahtis et al. [59] | 11 males, 3 females /62 ± 12 years/23 ± 8 days  | 14 ischemic/8 subcortical, 6 cortical/8 right, 6 left | (1) 2.0 mA bilateral tDCS         | IL 3.0 cm lateral to CZ | CL 3.0 cm lateral to CZ                        | 1 session à 15 min/pre post                            | TUG, POMA               |
| Manji et al. [53]  | 21 males, 9 females /63 ± 11 years/33 ± 9 days  | 17 ischemic, 13 hemorrhagic/na/na              | (1) 1.0 mA anodal+cathodal tDCS  | 35 cm anterior to CZ | inion                                         | 2 × 5 session à 20 min/pre, post1, post2              | 10MWT, FMA-LE, TUG, TCT, POMA                           |
| Sayes et al. [57]  | 17 males, 14 females /63 ± 9 years/42 ± 18 days | 26 ischemic, 5 hemorrhagic/7 subcortical, 24 cortical/14 right, 17 left | (1) 1.5 mA bilateral tDCS        | IL C3/C4               | CL C3/C4                                      | 16 sessions à 20 min/pre post                          | TT, RMI, TIS            |
| Andrade et al. [45] | 35 males, 25 females /69 ± 3 years/2.7 ± 0.5 months | na/na/na                                       | (1) 2.0 mA anodal tDCS           | IL C3/C4               | CL SO                                         | Parallel groups (15 + 15 + 15) + 10-20 EEG system    | FSST, OSI, BBS, FES-I, STS, 6MWT,                     |
| Kromjai et al. [50] | 14 males, 5 females /57 ± 3 years/32 ± 1,7 months | 19 ischemic/16 subcortical, 3 cortical/12 right, 7 left | (1) 2.0 mA bilateral tDCS        | IL M1                  | CL M1                                         | Crossover (19–19)/10–20 EEG system                    | TUG, STS, MVC (knee extensor)                          |
| Geroin et al. [49] | 14 males, 6 females /63 ± 7 years/26 ± 6 months | na/15 cortical, 5 subcortical/na               | (1) 1.5 mA anodal tDCS           | IL leg M1              | CL SO                                         | Parallel groups (10 + 10)/na                         | 6MWT, 10MWT, FAC, MI-LE, RMI, gait (cadence, symmetry, support time) |
| References                        | Participants number, gender, age/time since stroke | Stroke etiology/location/affected hemisphere | Stimulation technique, intensity | Electrodes positioning | Study design/electrodes positioning technique | Number and duration of sessions/evaluations schedule | Outcomes                                                                 |
|----------------------------------|---------------------------------------------------|---------------------------------------------|---------------------------------|------------------------|-----------------------------------------------|--------------------------------------------------|--------------------------------------------------------------------------|
| Picelli et al. [56]              | 22 males, 8 females/63 ± 8 years/4.7 ± 2.8 years | 30 ischemic/11 subcortical, 19 cortical/na  | (1) 2.0 mA anodal tDCS + sham tDCS | IL C3/C4               | CL SO                                          | Parallel groups (10 + 10)/10–20 EEG system          | 10 sessions à 40 min/pre, post, two-week, and four-week follow-up 6MWT, gait (cadence, support time) |
|                                 |                                                   |                                             | (2) 2.5 mA cathodal tDCS + sham tDCS | NA shoulder            | 10Th                                          |                                                   |                                                                         |
|                                 |                                                   |                                             | (3) 2.0 mA anodal tDCS + 2.5 mA cortical tDCS | IL C3/C4 + NA shoulder | CL SO + 10Th                                 |                                                   |                                                                         |
| Picelli et al. [55]              | 13 males, 7 females/s/63 ± 12 years/5.0 ± 3.7 years | 20 ischemic/7 subcortical, 13 cortical/na  | (1) 2.0 mA cathodal tDCS        | A buccinator muscle  | CL O1/O2                                       | Parallel groups (10 + 10)/10–20 EEG system          | 10 sessions à 20 min/pre, post, two-week, and four-week follow-up 6MWT, FAC, MI-LE, MAS, gait (cadence, support time) |
|                                 |                                                   |                                             | (2) 2.0 mA anodal tDCS          | IL leg M1              | CL SO                                         |                                                   |                                                                         |
| Picelli et al. [54]              | 21 males, 19 females/s/65 ± 10 years/5.3 ± 3.7 years | 40 ischemic, 14 subcortical, 26 cortical/na | (1) 2.0 mA cathodal tDCS        | A buccinator muscle  | CL O1/O2                                       | Parallel groups (20 + 20)/10–20 EEG system          | 10 sessions à 20 min/pre, post, two-week, and four-week follow-up 6MWT, gait (cadence, support time) |
|                                 |                                                   |                                             | (2) 2.0 mA cathodal tDCS        | NA buccinator muscle   | IL O1/O2                                       |                                                   |                                                                         |
| Madhavan et al. [52]             | 30 males, 10 females/60 ± 9 years/5.8 ± 4.6 years | 25 ischemic, 14 hemorrhagic, 1 na/na/na    | (1) 2.0 mA anodal tDCS         | IL leg M1              | CL SO                                         | Parallel groups (20 + 20)/10–20 EEG system          | 12 sessions à 15 min/pre, post, three months follow-up 10MWT, 6MWT, BBS, TUG, mini-BEST, FMA-LE, ABC |
|                                 |                                                   |                                             | (2) sham tDCS                  |                        |                                               |                                                   |                                                                         |
| Koganemaru et al. [51]           | 8 males, 3 females/66 ± 4 years/6.2 ± 2.6 years | 4 ischemic, 7 hemorrhagic/11 subcortical/na | (1) 0.0–20 mA bilateral tACS   | IL tibialis anterior M1 | CL 3.0 cm lateral and 3.0 cm rostral to inion | Crossover (11–11)/TMS                               | 2 × 1 session à 20 min/pre1, post1, pre2, post2 10MWT, VAS, MAS |
|                                 |                                                   |                                             | (2) sham tACS                  |                        |                                               | Crossover (8–8)/TMS                                | 2 × 5 sessions à 20 min/pre1, post1, one-week follow-up1, pre2, post2, post2, one-week follow-up 2 |
|                                 |                                                   |                                             |                                |                        |                                               |                                                   |                                                                         |
| Zandvliet et al. [60]            | 12 males, 3 females/57 ± 10 years/9 ± 1.2 years  | 11 ischemic, 4 hemorrhagic/2 subcortical, 13 cortical/9 right, 6 left | (1) 1.5 mA anodal tDCS        | IL 3.0 cm lateral to inion | NA buccinator muscle                           | Crossover (15–15–15)/10–20 EEG system              | 3 × 1 session à 20 min/pre1, post1, pre2, post3, eyes open, eyes closed and tandem stance (center of pressure amplitude, velocity, range) |
|                                 |                                                   |                                             | (2) 1.5 mA anodal tDCS          | CL 3.0 cm lateral to inion | A buccinator muscle                           |                                                   |                                                                         |
|                                 |                                                   |                                             | (3) sham tDCS                  |                        |                                               |                                                   |                                                                         |
| References          | Participants number, gender, age/time since stroke | Stroke etiology/ location/affected hemisphere | Stimulation technique, intensity | Electrodes positioning | Study design/ electrodes positioning technique | Number and duration of sessions/ evaluations schedule | Outcomes                                                                 |
|---------------------|-----------------------------------------------------|-----------------------------------------------|---------------------------------|------------------------|-----------------------------------------------|------------------------------------------------------|--------------------------------------------------------------------------|
| Seo et al. [58]     | 16 males, 5 females /62±9 years/9.5 ± 8.6 years     | 16 ischemic, 5 hemorrhagic/na/13 right, 8 left | (1) 2.0 mA anodal tDCS           | IL leg M1               | Parallel groups (11 + 10/10–20 EEG system)    | 10 sessions à 20 min/pre, post, four-week follow-up | 10MWT, 6MWT, FAC, BBS, FMA-LE, MRC (hip, knee, ankle)                 |
| Cattagni et al. [47]| 19 males, 5 females /57±13 years/10±7 years         | 17 ischemic, 7 hemorrhagic/na/11 right, 11 left | (1) 2.0 mA anodal tDCS           | IL leg M1               | Crossover (24–24)/10–20 EEG system            | 2 × 1 session à 30 min/pre, post                         | gait (speed, step length)                                               |
| Picelli et al. [55] | 13 males, 7 female s/63±12 years/5.0 ± 3.7 years    | 20 ischemic/7 subcortical, 13 cortical/na     | (1) 2.0 mA cathodal tDCS         | A buccinator muscle     | Parallel groups (10 + 10)/10–20 EEG system    | 10 sessions à 20 min/pre, two-week, and four-week follow-up | 6MWT, FAC, Mi-LE, MAS, gait (cadence, support time)                  |
| Picelli et al. [54] | 21 males, 19 female s/65±10 years/5.3 ± 3.7 years   | 40 ischemic, 14 subcortical, 26 cortical/na   | (1) 2.0 mA cathodal tDCS         | A buccinator muscle     | Parallel groups (20 + 20)/10–20 EEG system    | 10 sessions à 20 min/pre, two two-week, and four-week follow-up | 6MWT, gait (cadence, support time)                                   |
| Madhavan et al. [52]| 30 males, 10 females /60±9 years/5.8 ± 4.6 years    | 25 ischemic, 14 hemorrhagic, 1 na/na/na        | (1) 2.0 mA anodal tDCS           | IL leg M1               | Parallel groups (20 + 20)/10–20 EEG system    | 12 sessions à 15 min/pre, three months follow-up         | 10MWT, 6MWT, BBS, TUG, mini-BEST, FMA-LE, ABC                        |
| Koganemaru et al. [51]| 8 males, 3 females/66±4 years/6.2 ± 2.6 years       | 4 ischemic/7 hemorrhagic/11 subcortical/na    | (1) 0.0–20 mA bilateral tACS     | IL tibialis anterior M1 | Crossover (11–11)/TMS                         | 2 × 1 session à 20 min/pre1, post1, pre2, post2         | 10MWT, VAS, MAS                                                        |
| Zandvliet et al. [60] | 12 males, 3 females /57±10 years/9±1.2 years         | 11 ischemic, 4 hemorrhagic/2 subcortical, 13 cortical/9 right, 6 left | (1) 1.5 mA anodal tDCS           | IL 3.0 cm lateral to inion                     | Crossover (15–15–15)/10–20 EEG system                 | 3 × 1 session à 20 min/pre1, post1, pre2, post2, pre3, post3         | eyes open, eyes closed and tandem stance (center of pressure amplitude, velocity, range) |


| References          | Participants | Stroke etiology/ location/affected hemisphere | Stimulation technique, intensity | Electrodes positioning | Study design/ electrodes positioning technique | Number and duration of sessions/ evaluations schedule | Outcomes                      |
|---------------------|--------------|-----------------------------------------------|---------------------------------|------------------------|-----------------------------------------------|------------------------------------------------------|-------------------------------|
| Seo et al. [58]     | 16 males, 5 females /62±9 years/9.5±8.6 years | 16 ischemic, 5 hemorrhagic/na/13 right, 8 left | (1) 2.0 mA anodal tDCS          | IL leg M1               | Parallel groups (11+10)/10–20 EEG system            | 10 sessions à 20 min/pre, post, four-week follow-up | 10MWT, 6MWT, FAC, BBS, FES-I (hip, knee, ankle) |
| Cattagni et al. [47]| 19 males, 5 females /57±13 years/10±7 years  | 17 ischemic, 7 hemorrhagic/na/13 right, 11 left | (1) 2.0 mA anodal tDCS          | IL leg M1               | Crossover (24–24)/10–20 EEG system                  | 2 × 1 session à 30 min/pre, post gait (speed, step length) |                                |

BBS Berg Balance Scale, CL contralesional, cm centimeter, EEG electroencephalography, FES-I Falls Efficacy Scale—International, FMA-LE Fugl-Meyer Assessment—Lower Extremity, FSS T Four Square Step Test, IL ipsilesional, mA milliampere, min minute, MI-LE Motoricity Index—Lower Extremity, MRI magnetic resonance imaging, MVC maximal voluntary contraction, M1 primary motor cortex, na not available/not applicable, OSI Overall Stability Index, POMA Performance Oriented Mobility Assessment, RMI Rivermead Mobility Index, SO supra-orbital, STS Sit to Stand Test, tADS transcranial alternating current stimulation, TCT Trunk Control Test, tDCS transcranial direct current stimulation, Th10 the tenth thoracic vertebra, TIS Trunk Impairment Scale, TMS transcranial magnetic stimulation, tsDCS trans-spinal direct current stimulation, TT Tinetti Test, TUG Timed Up and Go Test, 6MWT Six-Minute Walk Test, 10MWT 10-Meter Walking Test, A affected, ABC Activities-Specific Balance Confidence Scale, BBS Berg Balance Scale, CL contralesional, cm centimeter, EEG electroencephalography, FAC Functional Ambulation Categories, FMA-LE Fugl-Meyer Assessment—Lower Extremity, mA milliampere, MAS Modified Ashworth Scale, min minute, MI-LE Motoricity Index—Lower Extremity, IL ipsilesional, mini-BEST mini-Balance-Evaluation-System-Test, MRC Medical Research Council, M1 primary motor cortex, NA non-affected, SO supra-orbital, VAS Visual Analog Scale,
## Table 2  Methodological quality of the included studies—assessed with the 11-item PEDro scale

| PEDro scale items | tDCS/tACS/tsDCS studies | rTMS studies |
|-------------------|--------------------------|--------------|
| Eligibility criteria specified | + | + |
| Random allocation | 1 1 1 1 1 1 1 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Concealed allocation | 1 0 0 0 1 0 0 1 0 0 1 0 0 1 1 1 0 0 1 0 0 1 1 1 0 |
| Comparable baseline | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Subject blinding | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Therapist blinding | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Assessor blinding | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Intention-to-treat analysis | 0 0 1 0 0 0 0 1 0 1 0 1 0 1 1 0 0 1 0 0 1 0 0 1 0 | 0 0 1 0 1 0 1 1 1 0 1 1 1 0 1 1 1 0 1 1 1 0 1 1 1 0 1 |
| Less than 15% dropouts | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| Between-group comparison | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |

*Note: The table shows the methodological quality assessment of the included studies using the PEDro scale. The scores are indicated with + for yes and 0 for no. The studies listed are: Bornheim et al. [46], Chang et al. [48], Tahtis et al. [59], Manji et al. [53], Sayes et al. [57], Andrade et al. [45], Klomjai et al. [50], Geroin et al. [49], Madhavan et al. [52], Picelli et al. [56], Picelli et al. [54], Picelli et al. [55], Madhavan et al. [52], Koganemaru et al. [51], Zandvliet et al. [60], Seo et al. [58], Cattagni et al. [47], Sasaki et al. [67], Kim et al. [63], Huang et al. [62], Koch et al. [64], Chiefflo et al. [61], Wang et al. [20], Rastgoo et al. [66], Wang et al. [68].*
Table 2 (continued)

| PEDro scale items | tDCS/tACS/tsDCS studies | rTMS studies |
|-------------------|-------------------------|--------------|
| Bornheim et al. [46] | et al. [48] | et al. [49] | 1 |
| Chang et al. [48] | et al. [50] | et al. [51] | 1 |
| Tahtis et al. [59] | et al. [52] | et al. [53] | 1 |
| Manji et al. [53] | et al. [54] | et al. [55] | 1 |
| Sayes et al. [57] | et al. [56] | et al. [57] | 1 |
| Andrade et al. [45] | et al. [58] | et al. [59] | 1 |
| Klomjai et al. [50] | et al. [60] | et al. [61] | 1 |
| Geroin et al. [49] | et al. [62] | et al. [63] | 1 |
| Madhavan et al. [52] | et al. [64] | et al. [65] | 1 |
| Picelli et al. [54] | et al. [66] | et al. [56] | 1 |
| Picelli et al. [55] | et al. [68] | et al. [57] | 1 |

Point estimates and variability

PEDro score total (0–10)

| 7 | 8 | 8 | 9 | 8 | 7 | 9 | 6 | 9 | 9 | 9 | 9 | 5 | 7 | 9 | 7 | 9 | 9 | 8 | 8 | 9 | 8 | 6 | 7 |

PEDro: Physiotherapy Evidence Database, rTMS: repetitive transcranial magnetic stimulation, tACS: transcranial alternating current stimulation, tDCS: transcranial direct current stimulation, tsDCS: trans-spinal direct current stimulation.
ipsilesional and the contralesional hemisphere) [61, 65, 67]. One study tested high-frequency rTMS over the ipsilesional hemisphere [68]. One trial applied iTBS over the contralesional hemisphere [64].

**Effectiveness—active stimulation versus sham:** In total, the post-interventional data indicate a small effect of rTMS on the observed parameters (Fig. 2). Despite a high homogeneity of detected effects, stimulation protocol dependent differences were found. High-frequency rTMS over the ipsilesional hemisphere induces large effects. ITBS over contralesional hemisphere evokes middle-sized effects. ITBS over contralesional hemisphere results in a small effect. No effect is induced by low-frequency rTMS over the contralesional hemisphere.

**Discussion**

Our data demonstrates that non-invasive neuromodulation is an effective way for improving gait, balance and/or lower limb motor function in stroke patients. This observation is supported by previous reviews and meta-analyses [69–72] that also demonstrate outcome-dependent effects [71, 72]. A recent meta-analysis revealed for example, significant effects of tDCS on functional ambulation category, Rivermead Mobility, and timed up and go test, but not on walking speed, 6-min walking distance, Tinetti test and Berg Balance Scale [71]. Similarly, another meta-analysis detected no relevant effects of rTMS on the Berg Balance Scale, while the remaining outcome measures for gait, balance and lower limb motor function were significantly influenced by the treatment [72]. In contrast to these studies, our meta-analysis focuses on the effectiveness in relation to the stimulation technique, protocol, hemisphere, area, duration and time since stroke (Table 4).
Our data shows that the superior effect of tDCS/tACS/tsDCS is primarily based on very high effects detected in various techniques on the central nervous system and its plasticity. It is an open question whether and to what extent the detected differences reflect the differential impact of various techniques on the central nervous system and behavior, and which role other factors (such as patients characteristics, study design, and “outlier” etc.) may play. Our data shows that the superior effect of tDCS/tACS/tsDCS is primarily based on very high effects detected in a single study, which performed three different experiments in large cohorts [45]. Therefore, the results of this trials. Similar observations were not made previously [69, 70]. It is an open question whether and to what extent the detected differences reflect the differential impact of various techniques on the central nervous system and behavior, and which role other factors (such as patients characteristics, study design, and "outlier" etc.) may play. Our data shows that the superior effect of tDCS/tACS/tsDCS is primarily based on very high effects detected in a single study, which performed three different experiments in large cohorts [45]. Therefore, the results of this

**Stimulation technique-dependent effects**

Both, the post-interventional data, and the follow-up data indicate that tDCS/tACS/tsDCS is superior to rTMS in supporting gait, balance, and lower limb function in stroke patients. These findings contrast with observations made in previous meta-analyses, that indicate superior effects of rTMS (in comparison to tDCS) on balance and postural control [70], or on hemi-spatial neglect [36] in this cohort. Furthermore, our data detects heterogeneity of tDCS/tACS/tsDCS effects that did not exist in rTMS.
study strongly influence the overall outcome of the meta-analysis. The reason for the superior stimulation effect- 
viness in this specific study may be the high amount of 
therapy applied in an early phase after the incident (a 
more detailed discussion of this topic is included below). 

From a global perspective, our data reveals that available 
rTMS and tDCS/tACS/tsDCS trials differ significantly 
with respect to the time since stroke, amount of inter-
vention, scheduling of the evaluations and study design 
(Figs. 2, 3, 4). While an average of 143 min (and 232 min 
in follow up data) of active treatment was applied in 
tDCS/tACS/tsDCS trials, an average of only 132 min 
(and 172 min in follow up data) was administrated in 
rTMS studies. This discrepancy may explain lower effect 
sized induced by rTMS interventions. The present data 
indicates that the behavioral changes are crucially deter-
mined by the amount of stimulation applied. Studies with 
relevant effects (according to Cohen's effect size defini-
tion) applied on average 169 min (and 232 min on follow 
up) of stimulation, while studies without relevant effects 
applied 99 min (and 172 min on follow up) of stimulation 
only. Furthermore, the data shows stimulation technique-dependent differences that indicate that only 
a higher amount of tDCS/tACS/tsDCS but not of rTMS 
induces larger behavioral changes. tDCS/tACS/tsDCS studies 
with statistically relevant effects applied on average 
186 min (and 241 min on follow up) of stimulation. 
tDCS/tACS/tsDCS trials without relevant effects applied 
89 min (and 196 min on follow up) of stimulation only. 
By contrast, rTMS was applied for 132 min (172 min 
on follow up) and 134 min (no follow up data) in studies 
with and without relevant effects, respectively. Our 
observations do not confirm a previous meta-analysis 
that suggested that a larger number of rTMS sessions 
corresponded to more benefits for balance and postural 
control in stroke patients [70]. Furthermore, significantly 
different stroke chronicity was detected in rTMS trials 
as compared with tDCS/tACS/tsDCS studies. While on 
average 43 months (and 35 months in follow up data) 
pass by since stroke in tDCS/tACS/tsDCS trials, there 
were only 13 months (18 months in follow up data) since 
the incident in rTMS studies. This is surprising, since 
the available data indicates that the chronicity of stroke 
correlates with less motor recovery, and reduced ther-
apy-induced benefits. A constraint-induced movement 
therapy applied over two weeks, for example, induces a 
greater improvement of motor function of the affected 
hand in patients who are less than 9 months post-stroke 
than in patients who are more than 12 months post-
stroke [73]. Moreover, the follow-up duration is on aver-
age ten weeks in tDCS/tACS/tsDCS studies and only four 
weeks in rTMS trials. Furthermore, no rTMS trial per-
forms a direct comparison of different protocols, in con-
trast to tDCS/tACS/tsDCS studies.

Stimulation protocol-dependent effects
Both the post-interventional and the follow-up data demon-
strate that different stimulation protocols induce dif-
ferential effects on gait, balance, and lower limb motor 
function in stroke patients. In sum, bilateral stimulation 
is superior to unilateral protocols. Similarly, a previous 
meta-analysis demonstrates the superior effectiveness of 
bilateral tDCS/rTMS for improving visuo-spatial abil-
ity following stroke [39]. However, a current meta-anal-
ysis indicates that bilateral tDCS is inferior to unilateral 
tDCS for improving motor learning in stroke patients 
[74]. Interestingly, our data shows that only tDCS stud-
ies apply the bilateral stimulation in accordance with the 
assumed maladaptive role of the contralesional hemo-
sphere [37]—with an anode over the ipsilesional and a 
cathode over the contralesional hemisphere [45, 50, 51, 
57, 59]. In contrast, bilateral rTMS studies applied high-
frequency protocols over either hemisphere [61, 65, 67].
| References       | Participants number, gender, age/time since stroke | Stroke etiology/location, affected hemisphere | Stimulation parameters, stimulated area | Study design/rTMS positioning technique | Number and duration of sessions/evaluations schedule | Outcomes                          |
|------------------|----------------------------------------------------|-----------------------------------------------|----------------------------------------|----------------------------------------|------------------------------------------------|-----------------------------------|
| Sasaki et al. [67] | 13 males, 8 females/11 ± 7 days                    | 11 ischemic, 10 hemorrhagic/na/9 right, 12 left | (1) 10 Hz bilateral rTMS (1000 pulses, 90% rMT, double-cone coil) over leg M1 (2) sham rTMS (sham coil) | Parallel groups (11 + 10)/TMS | 5 sessions à 10 min/pre, post | BRS                               |
| Kim et al. [63]   | 17 males, 15 females/16 ± 9 days                   | 32 ischemic/9 subcortical, 23 brain stem/na   | (1) 1 Hz unilateral rTMS (900 pulses, 100% rMT, figure-of-eight coil) over contralesional cerebellum (2) sham rTMS (perpendicular coil positioning) | Parallel groups (22 + 10/10–20 EEG system) | 5 sessions à 15 min/pre, one month follow up | 10MWT, BBS                       |
| Huang et al. [62] | 23 males, 15 females/29 ± 22 days                 | 25 ischemic, 13 hemorrhagic/28 subcortical, 10 cortical/17 right, 21 left | (1) 1 Hz unilateral rTMS (900 pulses, 120% rMT, double-cone coil) over contralesional musculus rectus femoris M1 (2) sham rTMS (sham coil) | Parallel groups (18 + 20)/TMS | 13 sessions à 15 min/pre, post, three months follow up | TUG, PASS, FMA-LE               |
| Lin et al. [65]   | 17 males, 3 females/13 ± 7 months                  | 16 ischemic, 4 hemorrhagic/9 right, 11 left   | (1) iTBS bilateral (1200 pulses, 100% rMT, figure-of-eight coil) over musculus rectus femoris M1 (2) sham iTBS (sham coil) | Parallel groups (10 + 10)/TMS | 10 sessions à 7 min/pre, post | BBS, TUG, 10MWT, FMA-LE           |
| Koch et al. [64]  | 21 males, 13 females/13 ± 17 months                | 34 ischemic/17 subcortical, 17 cortical/20 right, 14 left | (1) iTBS unilateral (1200 pulses, 80% aMT, figure-of-eight coil) over contralesional lateral cerebellum (2) sham iTBS (na) | Parallel groups (17 + 17)/neuronavigation | 15 sessions à 10 min/pre, three weeks follow up | BBS                               |
| Chiefflo et al. [61] | na/61 ± 10 years/20 ± 7 months                     | 5 ischemic, 4 hemorrhagic/9 subcortical/5 right, 4 left | (1) 20 Hz bilateral rTMS (1500 pulses, 90% rMT, H12-coil) over leg M1 (2) sham rTMS (sham coil) | Crossover (9–9)/TMS | 2 x 11 sessions à 30 min/pre, four weeks follow up | FMA-LE, 10MWT, 6MWT             |
| Wang et al. [20]  | 15 males, 9 females/23 ± 14 months                 | na/na/14 right, 10 left                       | (1) 1 Hz rTMS unilateral (600 pulses, 90% rMT, figure-of-eight coil) over contralesional musculus rectus femoris M1 (2) sham rTMS (perpendicular coil positioning) | Parallel groups (14 + 14)/TMS | 10 sessions à 10 min/pre, post | FMA-LE, gait analysis (speed, cadence, step length, support time, asymmetry) |
| References          | Participants number, gender, age/time since stroke | Stroke etiology/location/affected hemisphere | Stimulation parameters, stimulated area | Study design/rTMS positioning technique | Number and duration of sessions/evaluations schedule | Outcomes                                                                 |
|---------------------|----------------------------------------------------|---------------------------------------------|----------------------------------------|----------------------------------------|------------------------------------------------|--------------------------------------------------------------------------|
| Rastgoo et al. [66] | 16 males, 4 females/29±19 months                  | 15 ischemic, 5 hemorrhagic/15 subcortical, 5 cortical/13 right, 7 left | (1) 1 Hz rTMS unilateral (1000 pulses, 90% rMT, figure-of-eight coil) over contralesional leg M1 | Crossover (10–10)/TMS                  | 2 × 5 sessions à 17 min/pre1, post1, one week follow up1, pre2, post2, one week follow up2 | TUG, FMA-LE                                                              |
|                     |                                                    |                                             | (2) sham rTMS (sham coil)               |                                        |                                                                              |                                                                          |
| Wang et al. [68]    | 11 males, 3 females/29±20 months                  | 6 ischemic, 8 hemorrhagic/na/8 right, 6 left | (1) 5 Hz rTMS unilateral (900 pulses, 90% rMT, figure-of-eight coil) over ipsilesional musculus tibialis anterior M1 | Parallel groups (8 + 6)/TMS            | 9 sessions à 15 min/pre, post, one month follow up                  | FMA-LE, gait analysis (speed, asymmetry)                                |
|                     |                                                    |                                             | (2) sham rTMS (perpendicular coil positioning) |                                        |                                                                              |                                                                          |

BBS Berg Balance Scale, BRS Brunstrom Recovery Stages, EEG electroencephalography, FMA-LE Fugl-Meyer Assessment—Lower Extremity, Hz hertz, iTBS Intermittent theta burst stimulation, M1 primary motor cortex, min minute, na not applicable/not available, PASS Postural Assessment Scale for Stroke Patients, rMT resting motor threshold, (r)TMS (repetitive) transcranial magnetic stimulation, TUG Timed Up and Go Test, 6MWT Six-Minute Walk Test, 10MWT 10-Meter Walking Test
Table 4  Cross-tabulation of postinterventional effect sizes for studies evaluating non-invasive brain stimulation for improving gait, balance and/or lower limb recovery in stroke patients

| Stimulation induced effects on gait, balance and/or lower limb recovery (number of studies) | tDCS/tACS | rTMS |
|---|---|---|
| | Small negative effect | No effect | Small positive effect | Medium positive effect | Large positive effect | Small negative effect | No effect | Small positive effect | Medium positive effect | Large positive effect |
| Overall | 1 | 7 | 5 | 2 | 4 | 1 | 3 | 1 | 2 | 2 |
| Stimulated hemisphere | | | | | | | | | | |
| Affected | | | | | | | | | |
| Non-affected | 1 | 5 | 2 | 1 | 1 | | | | |
| Bilateral | 2 | 2 | 2 | 1 | 2 | | | | |
| Centred | 1 | | | | | | | | |
| Spinal | | | | | | | | | |
| Stimulated areas | | | | | | | | | |
| M1 | 1 | 4 | 3 | 2 | 2 | 1 | 3 | 2 | 2 |
| CR | 2 | | | | | 1 | | 1 | |
| M1 + M1 | 1 | 2 | 1 | 1 | 2 | | | | |
| M1 + CR | | | | | | | | | |
| SMA + M1 | 1 | | | | | | | | |
| 10Th | | | | | | | | | |
| Stimulation protocol | | | | | | | | | |
| 1 Hz rTMS | | | | | | 1 | | |
| 5 Hz rTMS | | | | | | 1 | | |
| 10 Hz rTMS | | | | | | 1 | | |
| 20 Hz rTMS | | | | | | 1 | | |
| iTBS | | | | | | | 1 | | |
| 0.0–2.0 mA tACS | | | | | | 1 | | |
| 1.0 mA anodal tDCS | | | | | | 1 | | |
| 1.5 mA anodal tDCS | 1 | 2 | | | | | | |
| 2.0 mA anodal tDCS | 4 | 1 | | | | | | |
| 1.0 mA anodal + cathodal tDCS | | | 1 | | | | | |
| 1.5 mA anodal + cathodal tDCS | | | 1 | | | | | |
| 2.0 mA anodal + cathodal tDCS | 1 | 1 | 1 | | | | | |
| 2.0 mA cathodal tDCS | | | | 1 | | | | |
| 2.5 mA cathodal tDCS | | | | | | 1 | | |
| Stimulation duration | tDCS/tACS |                  |                  |                  | rTMS |                  |                  |                  |
|----------------------|----------|-----------------|-----------------|-----------------|------|-----------------|-----------------|-----------------|
|                      | Small    | No effect       | Small positive  | Medium positive | Large | Small            | No effect       | Small positive  |
| ≥ 99 min             | negative |                 | effect          | effect          | effect|                 |                 | effect          |
| 100–199 min          | 1        | 3               | 2               | 1               | 1    | 2               | 1               | 1               |
| 200–299 min          | 3        |                 | 2               | 1               | 1    | 1               | 1               | 2               |
| ≤ 300 min            | 1        | 1               |                 |                 | 1    |                 |                 |                 |
| Time since stroke    |          |                 |                 |                 |      |                 |                 |                 |
| 0.1–3.2 months       | 1        | 2               | 2               | 1               | 3    | 1               | 1               | 3               |
| 11.0–290 months      | 2        | 0               |                 |                 | 2    | 1               | 2               | 2               |
| 42.0–1 200 months    | 5        | 3               | 1               | 1               |      |                 |                 |                 |

No effect (d < 0.2), small effect (d = 0.2–0.49), medium effect (d = 0.5–0.79), large effect (d ≥ 0.8), CR cerebellum, Hz hertz, iTBS intermittent theta burst stimulation, mA milliampere, M1 primary motor cortex, rTMS repetitive transcranial magnetic stimulation, SMA supplementary motor area, tACS transcranial alternating current stimulation, tDCS transcranial direct current stimulation, tsDCS trans-spinal direct current stimulation, T10Th the tenth thoracic vertebra.
The unilateral application of non-invasive brain stimulation induces, other than bilateral protocols, less consistent results across rTMS and tDCS/tACS/tsDCA studies. On the one hand, cathodal tDCS over the contralesional hemisphere induces large effects on gait, balance, and lower limb motor function [45]. On the other hand, low-frequency rTMS over this hemisphere had no effects [20, 62, 63, 66]. Interestingly, “facilitatory” stimulation of the contralesional hemisphere (contrary to its assumed maladaptive role) shows significant benefits with iTBS [64], but not with anodal tDCS [60]. Similarly, “facilitatory” stimulation of the ipsilesional hemisphere seems to be effective when using high-frequency rTMS [68], but not anodal tDCS [45, 47–49, 52, 58, 60]. Furthermore, cathodal tDCS56 and the coupling of frontal anodal tDCS with parietal cathodal tDCS [53] did not induce relevant effects. Collectively, the data indicates that the application of “facilitatory” TMS protocols (high-frequency rTMS, iTBS) induces an improvement of the assessed parameters, regardless of the stimulated hemisphere. In contrast, the successful application of tDCS is strongly determined by the concept of interhemispheric competition after a stroke, as described above [27]. In this context, it must be pointed out that recent data challenges the traditional view of either “facilitatory” or “inhibitory” effects of specific rTMS and tDCS protocols. It has been shown that rTMS (1 Hz/5 Hz/15 Hz/20 Hz), iTBS/cTBS, anodal/cathodal tDCS may all lead to both increases and decreases the corticospinal excitability [27–30]. The direction and/or the amount of TMS- and tDCS-induced changes may be significantly determined by individual (sex, age, genetics, medication, pre-interventional MEP latency and amplitude, pre-interventional MT size), technical (stimulator type, neuro-navigation use, TMS pulse waveform) and methodological (target muscle and hemisphere, time after stimulation, time of day, behavioral context) factors [28, 75–77]. A better understanding of the impact of these variables on stimulation effects may optimize the therapeutic application of these methods. Therefore, future studies need to develop more individualized stimulation protocols in accordance with the current knowledge [28, 75–77].

Beside this, only a limited spectrum of stimulation intensities has been investigated in previous stroke studies [20, 45, 68]. The intensity of 2.5 mA has not been exceeded in any tDCS trial. The existing evidence indicates, however, that higher tDCS intensities (≥ 2.0 mA) are more effective than low-intensity stimulation (≤ 1.5 mA) (Fig. 4). Moreover, recent studies show that intensities up to 4 mA are safe, tolerable, and do not elicit any serious adverse effects [78, 79]. Therefore, tDCS intensities between 3.0 and 4.0 mA may have the potential to better support stroke recovery than present protocols. Another relevant issue is the individualization of tDCS intensity. In contrast to rTMS which is applied in every subject with a specific intensity that is determined relative to the individual motor threshold, tDCS is usually applied with the same, predefined intensity in all subjects. It is, therefore, an open question, whether individualization of tDCS intensity corresponding to the respective corticospinal excitability may increase stimulation effectiveness [80]. Regarding rTMS, reasonably good evidence exists for the effects of 1 Hz rTMS supporting walking, balance, and lower limb function in stroke cohorts. However, the available data indicates better effectiveness of 5–20 Hz rTMS protocols (Table 4). Thus, stimulation frequencies above 20 Hz may evoke even higher effects on motor recovery than present protocols. However, the risk of rTMS-induced seizures increases also with rTMS frequency. Therefore, the risk–benefit ration should be carefully considered [81]. Up to now, stimulation protocols with frequencies of up to 50 Hz were successfully applied within the framework of neurorehabilitation [82]. Furthermore, most trials that were included in our meta-analysis applied rTMS with 90% of rMT. Only four studies applied another intensity (80% of active motor threshold [64], 100% [63, 65] and 120% [62] of resting motor threshold), and the available evidence did not show a superiority of any protocol.

A new way of tailoring TMS protocols is to consider the brain state of synchronized neuronal populations in the EEG at the time of stimulation. These EEG-triggered approaches may be informed by a series of post-hoc analyses of EEG features at the time of randomly applied TMS. In healthy subjects, stimulation effects on corticospinal excitability were less variable when the stimuli occurred at the optimum phase of beta frequency oscillations [83]. Along the same lines, the stimulation effects increased in both the resting and active motor system, when considering the oscillatory power of the beta-frequency band [84, 85]. Specifically, in both the resting brain85 and during voluntarily modulation [85], high and low beta-band activity decreased and increased corticospinal excitability, respectively. In addition, stimulation effects were modulated in a phase-dependent way along the oscillatory beta cycle and peaked with a diagonal shift of the highest stimulation response along the rising phase of the oscillatory cycle with increasing frequency [84]. Importantly, this phase-modulation was critically dependent on the precise temporal occurrence of the stimuli at a specific phase of the respective beta oscillatory cycle [86]. However, this high temporal precision could not be achieved in the past with EEG-triggered TMS approaches due to latencies between measurement and stimulation. Therefore, previous studies have applied EEG-controlled TMS on the basis of features that
necessitated less temporal precision such as high and low oscillatory power levels in the beta band (16–22 Hz) [31, 32], and positive and negative peaks of the slow (<1 Hz) [87] or alpha (8–12 Hz) [88, 89] oscillatory cycle. While beta power-dependent TMS induced robust increases of corticospinal excitability [31, 32], alpha peak-dependent findings were less consistent. Specifically, the alpha peak-dependent observations in a preselected group of participants with intrinsically high sensorimotor alpha power [88] could not be replicated, when the same approach of targeting the positive and negative peaks of the alpha cycle was applied in non-selected individuals [89]. Novel EEG-triggered approaches with integrated recording and stimulation devices and higher temporal precision may allow to repetitively target specific phases of higher frequency bands such as the oscillatory beta-band that have determined corticospinal excitability in previous post-hoc studies [83–86]. Moreover, such EEG-triggered approaches need also to be investigated in patient populations, e.g., following stroke [90], to explore their clinical utility under pathophysiological conditions.

**Stimulated area-dependent effect**

It is an open question, how much influence the stimulation location has on gait, balance, and lower limb motor function after stroke. Most analyzed studies stimulated the primary motor cortex. Beside this, cerebellum, supplementary motor area and spinal cord were also targeted in a few trials. However, a direct comparison of these areas is difficult because of numerous additional variables, such as different stimulation protocols, stimulated hemispheres, patient cohorts etc. Future studies need to create larger evidence for the application of non-invasive stimulation of areas other than the primary motor cortex. The cerebellum is a highly promising candidate in this regard. The available data indicates that the cerebellum is, similar to M1, critically involved in motor learning, but the mechanisms underlying cerebellar stimulation differ from those related to M1 stimulation. Specifically, the cerebellum is more linked to predictions about the consequences of movement than to direct motor commands [91]. Moreover, “cerebellar inhibition” (i.e., the inhibitory tone of the cerebellum over M1 via the thalamus) seems to play a key role during error-based motor learning, which is differently involved during early and late skill learning [91]. A current experiment demonstrates that preconditioning cerebellar stimulation improves not only the performance during the subsequent learning phase of visuo-motor adaptation tasks, but also induces a sustained improvement in the re-adaptation of the recently learned skill [92]. This observation is important for neurorehabilitation. It is an open question, however, whether and to which extend a stroke-induced damage of cortical motor areas may be compensated by cerebellar structures. In general, “cerebellar reserve” refers to the capacity of this area to compensate for tissue damage or loss of function following different etiologies [93]. Thus, it is plausible that cerebral stimulation may be a good alternative for patients that suffer from extensive cortical damage. Furthermore, it is conceivable that other cortical (such as the inferior parietal and frontal cortex) and subcortical (such as basal ganglia, thalamus, and hippocampus) regions and the brainstem are suitable for the application of non-invasive stimulation techniques for supporting gait and balance recovery [9, 13, 14]. Furthermore, the development of innovative technical devices enables modulating brain regions that could insufficiently be targeted by conventional stimulation equipment [26, 94, 95]. Double-cone coils, for example, are larger version of the standard figure-8 coils, and have two circular windings angled towards the subject’s head. Such double-cone coils are less focal but stimulate deeper brain areas than conventional figure-8 coils and may thus be beneficial for targeting the leg motor area, medial prefrontal cortex, cingulate, insula, and cerebellum [26, 94]. Similarly, other coil design (H, crown, stretched C-core, triple halo) have also the potential to modulate deeper brain areas than conventional figure-8 coils [94, 95], and their effectiveness in supporting gait, balance and lower limb function need to be investigated in future studies.

**Patient characteristic-dependent effects**

The studies included in our meta-analysis demonstrate an inconsistency of subjects regarding time since stroke, stroke etiology and lesion location. All these factors may hamper the interpretation of the results. Future studies should devote more attention to these important aspects. Studies investigating non-invasive brain stimulation for improving hand motor recovery could detect that lesion location may determine the effectiveness of the treatment [96, 97]. Fifteen sessions of 1 Hz rTMS over the contralateral primary motor cortex, for example, supported motor function of the affected hand only in patients with lesion of the dominant hemisphere. Patients with an injury of the non-dominant hemisphere did not profit from the intervention.98 Similarly, a single session of 10 Hz rTMS over the ipsilesional primary motor cortex significantly improved motor function of the affected upper limb in patients with a subcortical lesion. In contrast, no changes were detected in patients with cortical involvement [96]. Furthermore, a fMRI study detected different activation patterns during active movement of the affected lower limb in patients with subcortical and cortical stroke [98]. The data revealed similar activation patterns in patients with subcortical lesion and healthy controls with the recruitment of the contralateral
primary motor cortex, supplementary motor area, and bilateral somatosensory area. In contrast patients with cortical stroke and brainstem stroke showed reduced cortical recruitment [98]. Thus, it is conceivable that different stimulation protocols may be beneficial, depending on the lesion location.

**Strength and limitations**

To our knowledge, this is the first meta-analysis that compared the effectiveness of different non-invasive stimulation protocols in supporting gait, balance, and lower limb motor function in stroke subjects. An important strength is that we included and analyzed more studies than previously articles to this topic [69–72]. The main weakness is the inconsistency of analyzed studies with regard to the included patients (different time period since the incident, different stroke etiology and location), methodological approach (different numbers of intervention-sessions, different evaluation schedules), interventions (different stimulation protocols, different stimulation duration, different stimulated areas) and outcomes (more than twenty different assessments). Further weaknesses are the methodological limitations of the analyzed studies: (1) the absence of concealed allocation, (2) the absence of therapist blinding and (3) the absence of intention to treat analysis (Table 2). This may hamper the interpretation of the results.

**Conclusions**

This systematic review and meta-analysis show that certain types of non-invasive neuromodulation are effective in improving gait, balance, and lower limb motor function in stroke survivors. Available data indicates that (1) tDCS/tACS/tsDCS is more effective than rTMS, and that (2) bilateral stimulation is more effective than unilateral stimulation. However, more research is needed to maximize the effectiveness of existing protocols by optimizing stimulation dosage, intensity, and duration, by considering the brain state with EEG-triggered interventions, and by better characterizing the targeted stroke cohorts that may benefit.

**Abbreviations**

cTBS: Continuous theta burst stimulation; EEG: Electroencephalography; Hz: Hertz; iTBS: Intermittent theta burst stimulation; mA: Milliampere; MEP: Motor evoked potential; MR: Magnetic resonance imaging; ms: Millisecond; M1: Primary motor cortex; na: Not available/not applicable; PEDro: Physiotherapy Evidence Database; PRISMA: Preferred Reporting Items for Systematic Reviews and Meta-Analysis; TACS: Transcranial alternating current stimulation; tDCS: Transcranial direct current stimulation; tsDCS: Trans spinal direct current stimulation; rMT: Resting motor threshold; (r)TMS: (Repetitive) transcranial magnetic stimulation.

**Acknowledgements**

This work was supported by the German Federal Ministry of Education and Research (BMBF16SV8174, INERLINC). We acknowledge support by the Open Access Publishing Fund of the University of Tübingen.

**Author contributions**

JV and AG conceived and designed the study and performed the acquisition of the data. JV analyzed and interpreted the data and wrote the first version of the manuscript. AG contributed to data interpretation and reviewed the manuscript. Both authors read and approved the final manuscript.

**Funding**

Open Access funding enabled and organized by Projekt DEAL. This research received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors.

**Availability of data and materials**

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**

This article does not contain any studies with human participants or animals performed by any of the authors.

**Competing interests**

The authors declare no competing interests.

**Author details**

1Department of Sport Science, Bielefeld University, 33 501 Bielefeld, Germany.
2Institute for Neuromodulation and Neurotechnology, University Hospital and University of Tübingen, Tübingen, Germany.

Received: 15 February 2022   Accepted: 21 July 2022

Published online: 03 August 2022

**References**

1. Virani SS, Alonso A, Benjamin EJ, Bittencourt MS, Callaway CW, Carson AP, et al. Heart disease and stroke statistics—2020 update: a report from the American Heart Association. American Heart Association Council on Epidemiology and Prevention Statistics Committee and Stroke Statistics Subcommittee. Circulation. 2020;141:e139–596.
2. Jørgensen HS, Nakayama H, Raaschou HO, Olsen TS. Recovery of walking function in stroke patients: the Copenhagen Stroke Study. Arch Phys Med Rehabil. 1995;76:27–32.
3. Grau-Pellicer M, Chamarro-Lusar A, Medina-Casanovas J, Serdà Ferrer BC. Walking speed as a predictor of community mobility and quality of life after stroke. Top Stroke Rehabil. 2019;26:349–58.
4. Minet LR, Peterson E, von Koch L, Ytterberg C. Occurrence and predictors of falls in people with stroke: six-year prospective study. Stroke. 2015;46:2688–90.
5. Bohannon RW, Andrews AW, Smith MB. Rehabilitation goals of patients with hemiplegia. Int J Rehabil Res. 1988;11:181–3.
6. Pollock A, St George B, Fenton M, Finkins L. Top ten research priorities relating to life after stroke. Lancet Neurol. 2012;11:209.
7. Lang N, Siebner HR. Repetitive transcranial magnet stimulation. In: Siebner HR, Ziemann U, editors. The tRMS handbook. Heidelberg: Springer; 2007. p. 499–509.
8. Nitsche MA, Paulus W. Excitability changes induced in the human motor cortex by weak transcranial direct current stimulation. J Physiol. 2000;527:633–9.
with stroke: a systematic review and meta-analysis of randomized controlled trials. Clin Rehabil. 2019;33:1102–12.

37. Sawaki L, Butler AJ, Leng X, Wassenaar PA, Mohammad YM, Banton S, et al. Differential patterns of cortical reorganization following constraint-induced movement therapy during early and late period after stroke: a preliminary study. NeuroRehabilitation. 2014;35:415–26.

38. Halakoo S, Ehsani F, Masoudian N, Zoghli M, Jaberezhadi S. Does anodal trans-cranial direct current stimulation of the damaged primary motor cortex affects wrist flexor muscle spasticity and also activity of the wrist flexor and extensor muscles in patients with stroke? A randomized clinical trial. Neurourol Urodyn. 2021;42:2763–73.

39. Corp DT, Berzemicki HGK, Clark GM, Youssfj GF, Fried PJ, Jannati A, et al. ‘Big TMS Data Collaboration’ Large-scale analysis of interindividual variability in theta-burst stimulation data. Results from the ‘Big TMS Data Collaboration’. Brain Stimul. 2020; 13:1476–1488.

40. Corp DT, Berzemicki HGK, Clark GM, Youssufj GF, Fried PJ, Jannati A, et al. ‘Big TMS Data Collaboration’ Large-scale analysis of interindividual variability in single and paired-pulse TMS data. Clin Neurophysiol. 2021; 132:2639–2633.

41. Ridding MC, Ziemann U. Determinants of the induction of cortical plasticity by non-invasive brain stimulation in healthy subjects. J Physiol. 2010;588:2291–304.

42. Bickon M, Grossman P, Thomas C, Zannou AL, Jiang J, Adnan T, et al. Safety of transcranial direct current stimulation: evidence based update 2016. Brain Stimul. 2016;9:641–61.

43. Khademi F, Royster V, Gharabaghi A. Distinct beta-band oscillatory circuits state dependent stimulation: enhancing motor cortex excitability for neurorehabilitation. Brain Stimul. 2018;11:374–89.

44. Torrecillos F, Falato E, Pogosyan A, West T, Di Lazzaro V, Brown P. Motor cortex inputs at the optimum phase of beta cortical oscillations undergo more rapid and less variable corticospinal propagation. J Neurosci. 2012;32:243–53.

45. Bonnì S, Motta C, Pellicciari MC, Casula EP, Cinnera AM, Maiella M, et al. Safety study of 50 Hz repetitive transcranial magnetic stimulation over the primary motor cortex. Neuroimage Rep. 2020;40:369–81.

46. Corp DT, Bereznicki HGK, Clark GM, Youssefj GF, Fried PJ, Jannati A, et al. ‘Big TMS Data Collaboration’ Large-scale analysis of interindividual variability in single and paired-pulse TMS data. Clin Neurophysiol. 2021;132:2639–2633.

47. Ridding MC, Ziemann U. Determinants of the induction of cortical plasticity by non-invasive brain stimulation in healthy subjects. J Physiol. 2010;588:2291–304.

48. Bickon M, Grossman P, Thomas C, Zannou AL, Jiang J, Adnan T, et al. Safety of transcranial direct current stimulation: evidence based update 2016. Brain Stimul. 2016;9:641–61.

49. Khademi F, Royster V, Gharabaghi A. Distinct beta-band oscillatory circuits state dependent stimulation: enhancing motor cortex excitability for neurorehabilitation. Brain Stimul. 2018;11:374–89.

50. Torrecillos F, Falato E, Pogosyan A, West T, Di Lazzaro V, Brown P. Motor cortex inputs at the optimum phase of beta cortical oscillations undergo more rapid and less variable corticospinal propagation. J Neurosci. 2012;32:243–53.

51. Bonnì S, Motta C, Pellicciari MC, Casula EP, Cinnera AM, Maiella M, et al. Safety study of 50 Hz repetitive transcranial magnetic stimulation over the primary motor cortex. Neuroimage Rep. 2020;40:369–81.
93. Mitoma H, Buffo A, Gelfo F, Guell X, Fucà E, Kakei S, et al. Consensus paper. Cerebellar reserve: from cerebellar physiology to cerebellar disorders. Cerebellum. 2020;19:131–3.
94. Deng ZD, Lisanby SH, Peterchev AV. Coil design considerations for deep transcranial magnetic stimulation. Clin Neurophysiol. 2014;125:1202–12.
95. Rastogi P, Lee EG, Hadimani RL, Jiles DC. Transcranial magnetic stimulation: development of a novel deep brain coil—triple halo coil. IEEE Magn Lett. 2019;10:1–5.
96. Ameli M, Greffes C, Kemper F, Rehme AK, Karbe H, et al. Differential effects of high-frequency repetitive transcranial magnetic stimulation over ipsilesional primary motor cortex in cortical and subcortical middle cerebral artery stroke. Ann Neurol. 2009;66:298–309.
97. Lüdemann-Podubecká J, Bösl K, Theilig S, Wiederer R, Nowak DA. The effectiveness of 1 Hz rTMS over the primary motor area of the unaffected hemisphere to improve hand function after stroke depends on hemispheric dominance. Brain Stimul. 2015;8:823–30.
98. Luft AR, Forrester L, Macko RF, McCombe-Waller S, Whittall J, Villagra F, Hanley DF. Brain activation of lower extremity movement in chronically impaired stroke survivors. Neuroimage. 2005;26:184–94.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.