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Modulation of inhibitory systems to enhance motor rehabilitation: insights for the use of noninvasive brain stimulation


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Modulation of inhibitory systems to enhance motor rehabilitation: insights for the use of noninvasive brain stimulation


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Abstract

Motor impairment following stroke is a leading cause of disability in adults. Despite advances in motor rehabilitation techniques, many adult stroke survivors never approach full functional recovery. Intriguingly, children exhibit better rehabilitation outcomes when compared to adults suffering from comparable brain injuries, yet the reasons for this remain unclear. A common explanation is that neuroplasticity in adults is substantially limited following stroke, thus constraining the brain’s ability to reorganize in response to neurological insult. This explanation, however, does not suffice for there is much evidence suggesting that neuroplasticity in adults is not limited following stroke. We hypothesize that diminished functional recovery in adults is in part due to inhibitory neuronal interactions, such as transcallosal inhibition, that serve to optimize motor performance as the brain matures. Following stroke, these inhibitory interactions pose rigid barriers to recovery by inhibiting activity in the affected regions and hindering recruitment of compensatory pathways. In contrast, children exhibit better rehabilitation outcomes in part because they have not fully developed the inhibitory interactions that impede functional recovery in adults. We suggest that noninvasive brain stimulation can be used in the context of motor rehabilitation following stroke to reduce the effects of existing inhibitory connections, effectively returning the brain to a state that is more amenable to rehabilitation. We conclude by discussing further research to explore this hypothesis and its implications. Keywords: stroke, motor recovery, plasticity, rehabilitation, transcranial magnetic stimulation, transcranial direct current stimulation.

Introduction

Functional recovery following stroke poses a significant challenge. Current rehabilitation strategies adopt multidisciplinary approaches that include physical, occupational, cognitive, and other forms of therapy. The Bobath approach, for instance, is a widely utilized technique that aims to integrate postural control with task performance and to train coordinated sequences of movement for a given task (Graham, Eustace, Brock, Swain, & Irwin-Carruthers, 2009; Natarajan et al., 2008). Although this and other rehabilitation techniques have reduced morbidity and institutionalized care, functional outcomes remain limited (Desrosiers, Bourbonnais, Corriveau, Gosselin, & Bravo, 2005; Hendricks, van Limbeek, Geurts, & Zwarts, 2002; Indredavik, Bakke, Slordahl, Rokseth, & Haheim, 1999; Langhammer & Stanghellie, 2003). Indeed, motor impairment following stroke is a leading cause of disability in adults, with long term medical expenses resulting from stroke representing a majority of costs associated with this expensive disease (Demaerschalk, Hwang, & Leung, 2010). It is estimated that less than 15% of stroke patients exhibiting paralysis in the upper or lower extremities achieve complete motor recovery (Hendricks et al., 2002).

While functional improvement is limited in adults, interestingly, children suffering from comparable brain injuries exhibit substantially better rehabilitation outcomes. Following stroke, children often demonstrate
As mentioned above, one explanation for the relatively low functional recovery in adults is that neuroplasticity might be reduced in the brain following stroke. Here, we present evidence on the molecular, cellular, and systemic levels that suggests that neuroplasticity is not generally limited following stroke.

At the molecular level, studies have shown that following ischemic insult, there is a re-emergence of developmental proteins that are otherwise normally absent or present at very low concentrations (Cramer & Chopp, 2000; Stroemer et al., 1998). These proteins are associated with increased neuroplasticity and include differentiation factors (e.g., NeuroD), structural proteins (e.g., Nestin, MAP-2), growth associated proteins (e.g., GAP43, synaptophysin), as well as growth factors (e.g., BFGF, VEGF, BDNF) (Cramer & Chopp, 2000). Interestingly, these proteins are increased not only at the local site of injury, but in distant areas as well. Furthermore, while these proteins re-emerge within hours to days following the lesion, they remain effective for weeks and months after the initial insult (Cramer & Chopp, 2000).

At the cellular level, there is also convincing evidence of increased plasticity following injury. One study in adult rats demonstrated that the propensity for long-term potentiation is increased in perilesional areas after experimentally induced focal cortical infarction (Hagemann, Redecker, Neumann-Haefelin, Freund, & Witte, 1998). Similarly, unilateral injury to the sensorimotor cortex in adult rats has been shown to induce dramatic growth of neuronal dendrites in the contralateral homotopic cortex (Jones et al., 1996). Finally, neocortical injury has been shown to increase dendritic branching (Jones & Schallert, 1992) and synaptic density (Jones et al., 1996; Stroemer et al., 1998). These studies suggest that on both molecular and cellular levels, mechanisms geared toward promoting brain reorganization are enhanced, rather than inhibited, following injury.

At the systemic level, evidence suggests that dramatic cortical reorganization also takes place following stroke. A study in adult primates, for instance, found cortical remapping along the margins of the site of focal damage (Xerri, Merzenich, Peterson, & Jenkins, 1998). Furthermore, following focused rehabilitative training of skilled movements, the motor area representation of the impaired hand in primates have been shown to extend to areas formerly associated with shoulder and elbow movements (Nudo, Wise, SiFuentes, & Milliken, 1996). Consistent with the animal findings above, motor areas associated with finger movements in adult stroke patients have been shown to be remapped onto areas traditionally associated with face motor control (Weiller, Ramsay, Wise, Friston, & Frackowiak, 1993). Another study in similar patients found that recovery from stroke was associated with motor cortical reorganization including bihemispheric recruitment of premotor cortical areas (Seitz et al., 1998).

Studies in adult patients using functional magnetic resonance imaging (fMRI) and positron emission
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tomography (PET) provide robust evidence for cortical reorganization following stroke. A longitudinal fMRI study of 20 first-time adult stroke survivors, for instance, showed task-related changes in brain activation over the course of recovery (Ward, Brown, Thompson, & Frackowiak, 2003). An extensive review of PET and fMRI studies supports the conclusion that the damaged adult brain is indeed capable of reorganizing as a means of compensating for motor impairments (Calautti & Baron, 2003).

This cortical remapping has also been observed in healthy adults receiving low frequency (1Hz) repetitive transcranial magnetic stimulation (rTMS) delivered to the contralateral primary motor cortex (M1) (Lee et al., 2003). rTMS is a method of noninvasive brain stimulation that uses a large, rapidly changing magnetic field to induce electrical stimulating currents in the brain. At low frequency of stimulation, this procedure is presumed to decrease neuronal firing and excitability (Pascual-Leone et al., 2005; Pascual-Leone et al., 1998). Lee et al. (2003) found that subjects who performed simple motor tasks following rTMS of the motor cortex showed sustained motor function while having activation of secondary motor areas such as the rostral supplementary motor cortex, ipsilateral primary motor cortex, and dorsal premotor cortex (as detected by fMRI). They suggest that this rTMS-induced remapping may be similar to the acute compensatory plasticity of the motor system following stroke.

The above human and animal studies strongly support the notion that neuroplasticity is present in adults following stroke. While the studies do not provide direct comparisons to children, they do counter the claim that neuroplasticity is absent in adults and call into question whether differences in neuroplasticity alone can explain the poor functional recovery in adults following stroke.

Alternative neural pathways in stroke recovery

Alternative neural pathways, such as corticospinal projections from the ipsilesional dorsolateral premotor cortex (PMd) and supplementary motor area (SMA), appear to play important roles in stroke recovery. These projections are known to be less numerous and less effective at exciting spinal cord motor neurons when compared to projections from the primary motor cortex (Maier et al., 2002). However, several studies have shown that these alternative neural pathways are recruited in stroke recovery. For instance, there are reports of increased ipsilesional PMd activation associated with therapy-induced improvement in upper limb function (Johansen-Berg et al., 2002) and gait function (Miyai et al., 2003). Furthermore, chronic stroke patients exhibit simple task impairments when the ipsilesional PMd is disrupted by TMS (a finding not observed in healthy controls), demonstrating again the importance of this alternative neural pathway in motor recovery (Fridman et al., 2004).

While alternative neural pathways appear to play an important role in motor recovery following stroke, their role remains limited. Indeed, long-term motor recovery is more strongly associated with the extent and location of damage to direct corticospinal projections from the primary motor cortex, with greater preservation and activation of direct corticospinal tract following stroke predicting better functional outcomes in patients (Cruz Martinez, Tejada, & Diez Tejedor, 1999; Pennisi et al., 1999). Longitudinal studies have shown that enhanced motor recovery in adults is associated with cortical activation patterns more similar to those observed before stroke, suggesting that optimum recovery following stroke requires a shift from initial compensatory patterns of activation, using alternative neural pathways to pre-stroke patterns using the affected primary motor cortex (Jang et al., 2004; Jang et al., 2003; Ward et al., 2003).

In summary, while remapping to alternative neural pathways appears important for motor recovery following strokes, better results are achieved in patients who are able to eventually shift neural activity back to its pre-stroke state. It seems reasonable to infer that anatomical preservation of motor neural systems is important for such outcomes. Interestingly, this does not appear to be the case for children recovering from brain lesions. Studies have shown that even in extremes cases such as hemispherectomy, younger patients achieve significantly greater functional recovery than older subjects (Benecke et al., 1991; Gardner et al., 1955). Controlled animal studies support the conclusion that younger age predicts better outcomes as well. For instance, a study comparing motor outcomes in neonatal and adult hemispherectomized cats, found that the neonatal group relearned how to use their impaired limbs following one practice session while the adult group needed almost eight training sessions to acquire the same motor skill (Burgess, Villablanca, & Levine, 1986). A recent experiment directly comparing hemispherectomy in infant and adult monkeys similarly showed better functional outcomes in the younger animals (Burke, Zangenehpour, & Pito, 2010). As discussed above, these differences in outcomes across age cannot simply be due to an utter lack of neuroplasticity in older subjects, as neuroplasticity and recruitment of alternative neural pathways have been widely observed in adults. A deeper understanding of the effects of age on brain function, injury, and recovery may provide useful insight into these differences.

Maturation and inhibitory systems: age as an important determinant of stroke recovery

Maturation of the nervous system is a complex process involving elaborate formation, myelination, and elimination (pruning) of synaptic connections that starts in the uterus and extends into teenage years (Garvey...
With respect to motor function, maturation includes the development of handedness, loss of synkinesis (by age 10), and gradual increase in motor performance over time (Frye, Rotenberg, Ousley, & Pascual-Leone, 2008; Garvey & Mall, 2008; Mall et al., 2004). Interestingly, the combined results of these studies suggest that the relatively reduced inhibitory systems in children facilitate neuroplastic changes and recruitment of alternative neural pathways, making them more amenable to the neuroplastic processes necessary for recruiting alternative neural pathways, return to pre-stroke activation patterns, and achieving full functional recovery.

The observation that transcallosal inhibition hinders motor rehabilitation, may in part explain the better functional outcomes in pediatric patients as compared with adults following stroke and other forms of brain injury. As discussed above, children below the age of 10 have not fully developed the interhemispheric and intracortical inhibitory systems that impede full motor recovery (Heinen et al., 1998; Mall et al., 2004). Therefore, cortical excitability in their lesioned cortices will have greater cortical excitability than their adult counterparts, making them more amenable to the neuroplastic processes necessary for recruiting alternative neural pathways, returning to pre-stroke activation patterns, and achieving full functional recovery.

The suggestion that relatively low inhibitory drive in younger patients facilitates recovery may shed light on the remarkable functional outcomes observed in children following brain injury. As previously mentioned, research confirms that pediatric stroke survivors and neonatal hemispherectomy patients surpass their adult counterparts in outcome measures (Benecke et al., 1991; Gardner et al., 1955; Kim et al., 2009). One study comparing patients hemispherectomized before and after brain maturation found that TMS stimulation evoked faster and stronger ipsilateral compound muscle action potentials in patients that underwent hemispherectomy prior to brain maturation (Benecke et al., 1991). This study suggests that maturation of inhibitory systems not only renders alternative neural pathways less accessible in adulthood, but also less efficient, with slower and weaker responses than those observed in children. Conversely, the combined results of these studies suggest that the relatively reduced inhibitory systems in children facilitate neuroplastic changes and recruitment of compensatory pathways.

The above discussion suggests that diminished functional recovery in adults is in part due to inhibitory neuronal interactions, such as transcallosal inhibition, that serve to optimize motor performance as the brain matures. Following stroke, these inhibitory interactions pose rigid barriers to recovery by inhibiting activity in the affected regions and hindering recruitment of compensatory pathways. Building off of this
understanding, we next discuss noninvasive brain stimulation techniques that aim to modulate these inhibitory systems, thereby returning the brain to a state that is more amenable to rehabilitation.

**Using noninvasive brain stimulation to guide brain plasticity in stroke recovery**

Despite the many rehabilitative therapies that are currently employed, functional recovery following stroke remains fairly limited. Based on the above discussion, we propose the use of noninvasive brain stimulation, particularly transcranial magnetic stimulation (TMS) and transcranial direct current stimulation (tDCS), as techniques for modulating inhibitory systems that predominate in adults and for enhancing motor recovery following stroke.

The use of electro-magnetic brain stimulation has been investigated since the late 19th century (Walsh, Pascual-Leone, & Kosslyn, 2005). It was not until the mid 1980’s, however, that TMS was introduced, when Anthony Barker and colleagues solved the technical challenges involved in penetrating the scalp and skull with a sufficiently strong and quick magnetic field pulse (Barker, Jalinous, & Freeston, 1985). Whereas tDCS (discussed below) is considered to be a purely neuromodulatory technique, TMS is a neurostimulatory and neuromodulatory technique. TMS uses the principle of electromagnetic induction to produce localized, induced currents in the brain (Hallett, 2000). These currents can be of sufficient magnitude to depolarize neurons and stimulate action potentials, and when these currents are applied repetitively, as in repetitive transcranial magnetic stimulation (rTMS), they can also modulate cortical excitability beyond the time of stimulation, either decreasing or increasing general excitability (depending on the parameters of stimulation; Pascual-Leone et al., 1999).

**tDCS** is a less expensive and more portable technique that involves the transmission of low amplitude direct currents from sponge electrodes through the skull and to the underlying cortex. Although there is substantial shunting of current at the scalp, studies have shown that sufficient current penetrates the brain to modify the transmembrane resting potentials of cortical neurons and thereby influence their firing rate and general excitability in response to input (Miranda, Lomarev, & Hallett, 2006; Wagner et al., 2007). When tDCS is applied for a sufficient duration, it has after-effects on neuronal spontaneous activity as demonstrated in humans by shifts in cortical excitability that can be observed well beyond the period of stimulation (Nitsche & Paulus, 2001).

Below we suggest three ways in which noninvasive brain stimulation can be used to enhance patient care and motor recovery following stroke. First, we discuss the potential use of TMS as a diagnostic tool for recording neurophysiological measures and tracking improvement following stroke. We next discuss how rTMS and tDCS can be used to counteract maladaptive consequences of inhibitory systems and excite affected cortical areas, thereby facilitating motor recovery. Finally, we discuss how tDCS can be used to enhance synaptic plasticity and re-learning of motor skills following stroke.

**Transcranial magnetic stimulation as a diagnostic tool**

When applied to the motor cortex at a suitable intensity, TMS induces the depolarization of pyramidal cells, generating a descending corticospinal volley that depolarizes spinal motoneurons and results in the induction of a muscle contraction. The amplitude of the muscle contraction (motor evoked potential; MEP) is a measure of the activation of the motor cortex at a specific point in time. In addition, applied to motor and non-motor brain regions, TMS induces depolarization of cortical neurons and activates cortico-cortical and cortico-subcortical connections that induce specific changes in brain activity, which can be recorded using EEG. Such measures can provide insights into brain connectivity both within and across hemispheres. Importantly, the impact of TMS depends on the level of local cortical excitability and thus on the balance of intracortical facilitation and inhibition present in the brain. Therefore, neurophysiological measures obtained using TMS measures can provide unique insights regarding conditions in which there are imbalances in intracortical excitability, such as in stroke or brain damage.

Besides the assessment of motor evoked potential and motor threshold, which measure global corticospinal excitability levels, various measures of intra- and inter-cortical inhibition and facilitation are also possible with TMS. These include: (1) **short interval intracortical inhibition** (SICI) (Kujirai et al., 1993). It is assumed that SICI involves the inhibition of action potentials resulting from the second, supra-threshold pulse by the activation of low-threshold inhibitory circuits (via inhibitory post synaptic potentials; IPSP) generated by the first, sub-threshold pulse. It is widely believed that SICI is GABA$_{A_1}$-dependent since GABA$_{A_1}$ agonists increase SICI (Ziemann, Rothwell, & Ridding, 1996); (2) **long interval intracortical inhibition** (LICI), where both TMS pulses are delivered at supra-threshold intensities with an interstimulus interval of 50-200 ms. There is strong evidence that LICI is mediated by long-lasting GABA$_{A_1}$-dependent IPSPs and activation of pre-synaptic GABA$_{A_1}$ receptors on inhibitory interneurons (Werhahn, Kunesch, Noachtar, Benecke, & Classen, 1999); (3) **intracortical facilitation** (ICF), the amplitude of a test MEP can be enhanced if it is accompanied by a sub-threshold conditioning pulse applied 10-25 ms earlier (Kujirai et al., 1993). Glutamatergic interneurons at the level of M1 are likely to be involved in ICF since it is reduced by NMDA antagonists such as dextromethorphan; and (4) **transcallosal inhibition** in which the two motor cortices...
are stimulated with a delay of 10ms. The first pulse (the conditioning pulse) is applied over the primary motor cortex and the second pulse (the test pulse) is applied after a delay of 10ms in the contralateral primary motor cortex. It has been shown that the second pulse is associated with a significant inhibition in the MEP characteristics.

As discussed above, transcallosal inhibition begins developing in children 6 years of age (Heinen et al., 1998) and appears to have maladaptive effects in the context of motor recovery following stroke (Murase et al., 2004).

These neurophysiological parameters can give a relatively accurate estimate of the level of inhibition in the motor systems. TMS can therefore serve as a diagnostic tool to monitor the natural progression of stroke and measure the effects of therapies on recovery.

**rTMS and tDCS to modulate maladaptive inhibitory systems and increase motor recovery**

As discussed in this review, inhibitory interactions are strengthened during maturation, which limit the ability of the cortex to recruit alternative neural pathways. In fact, after stroke there are multifocal, bihemispheric changes in brain activity, which pose rigid barriers to recovery by inhibiting activity in the affected regions and hindering recruitment of compensatory pathways (Ward et al., 2003). An example here is the increased transcallosal inhibition that is observed in the acute phases following stroke (Murase et al., 2004).

Numerous investigations point to the conclusion that activation of the motor cortex is of paramount importance to promote motor recovery. Anatomically, it is the most common source of fibers for the pyramidal tract (Galea & Darian-Smith, 1994), which is key to generating rapid, distal, or fractionated movements (Brinkman & Kuypers, 1973). Studies in animals (Dijkhuizen et al., 2003) and humans (Marshall et al., 2000) have described a hemispheric shift in sensorimotor cortex activation from bilateral to stroke-affected hemisphere in association with post-stroke recovery, particularly among patients with subcortical stroke (Feydy et al., 2002; Theoret, Kobayash, Valero-Cabre, & Pascual-Leone, 2003).

Following stroke of the motor cortex, cortical excitability is relatively decreased as compared to the unaffected motor cortex (Shimizu et al., 2002) possibly due to a shift in interhemispheric interactions with increased transcallosal inhibition from the unaffected to the affected motor cortex (Murase et al., 2004). Similar shifts in interhemispheric interactions have been postulated across parietal cortices in strokes leading to neglect syndrome (Kinsbourne, 1977).

Modulation of interhemispheric competition using rTMS has been demonstrated in normal subjects. For example, low frequency rTMS can improve motor performance with the ipsilateral hand by releasing interhemispheric inhibition in the unstimulated hemisphere (Kobayashi, Hutchinson, Theoret, Schlaug, & Pascual-Leone, 2004). Similarly, low frequency rTMS of one parietal cortex improves ipsilateral attention by enhancing excitability in the unstimulated parietal cortex (Hilgetag et al., 2001). In the setting of a stroke, there might thus be desirable effects induced by either suppressing activity in the undamaged hemisphere or increasing activity in the perilesional cortex of the damaged hemisphere, thereby promoting a restoration of activity across bihemispheric neural networks and guidance of more adaptive plasticity (Fregni & Pascual-Leone, 2006; Hummel & Cohen, 2006).

Indeed, preliminary studies using rTMS to enhance motor recovery in stroke patients provide promising results. In a sham-controlled trial that included patients up to 12 months following stroke, Mansur and colleagues (2005) applied 0.5 Hz rTMS to the unaffected hemisphere for 10 minutes to suppress cortical activity and maladaptive transcallosal inhibition influencing the affected hemisphere. They found that the treatment significantly improved performance of simple and choice reaction time tests as well as the Purdue Pegboard test in the affected hand (Mansur et al., 2005). Extending these findings to children, a randomized study in 10 pediatric patients (>7 years old) who suffered from subcortical arterial ischaemic stroke found that contralesional inhibitory rTMS seemed to improve hand function in children with hemiparesis, with no serious adverse effects (Kirton et al., 2008).

Preliminary studies using tDCS to modulate motor cortex activity in stroke patients and healthy subjects have also returned favorable results. Fregni et al. (2005b) examined whether reduction of activity in the contralesional hemispheres of stroke patients by introducing cathodal (inhibitory) tDCS would result in improved motor performance due to decreased transcallosal inhibition. They found that cathodal stimulation of the contralesional hemisphere, as well as anodal (excitatory) stimulation of the ipsilesional hemisphere, significantly improved motor performance compared to sham tDCS (Fregni et al., 2005a). In a recent sham-controlled investigation of healthy subjects, Williams, Pascual-Leone, and Fregni (2010) combined bilateral motor cortex tDCS with contralateral hand restraint of the dominant hand. When comparing active- to sham- stimulation, they found a decrease in cortical excitability in the dominant hemisphere and a decrease in transcallosal inhibition from the dominant hemisphere to the non-dominant hemisphere. Moreover, this decrease in transcallosal inhibition was positively associated with motor performance enhancement in the non-dominant hand (Williams et al., 2010).

In sum, the above discussion suggests that both rTMS and tDCS may be useful tools for decreasing maladaptive interhemispheric inhibition, exciting lesioned cortex, and facilitating motor recovery following stroke.
Using tDCS to enhance the effects of cognitive-behavioral interventions

Following stroke, activation of neural networks that support re-learning of motor skills remains an important challenge. While physical therapy may be a valuable approach to achieving motor recovery, it is of limited efficacy when administered alone. Here we discuss how tDCS can be used to enhance the efficacy and efficiency of cognitive-behavioral interventions by promoting neuroplasticity and consolidation of newly formed synaptic connections.

The acquisition of new skills and behaviors is linked to changes in neuronal activity and excitability (Facchini, Muellbacher, Battaglia, Boroojerdi, & Hallett, 2002). This might reflect changes in synaptic strength, for example, through N-methyl-D-aspartate (NMDA) receptor-dependent long-term potentiation (LTP) (Rioult-Pedotti, Friedman, Hess, & Donoghue, 1998). Successful manipulation of cortical excitability to improve learning processes and cognitive training has been demonstrated in humans in neuropharmacological investigations (Butefisch et al., 2002), with TMS (Butefisch, Khurana, Kopylev, & Cohen, 2004), and with deafferentation of adjacent or contralateral body parts (Facchini et al., 2002; Floel et al., 2004). tDCS thus presents an interesting alternative because it is non-invasive, painless (as compared to TMS and deafferentation), and without serious side effects (as compared to neuropharmacological agents). In addition, tDCS has an important theoretical advantage as it directly modifies spontaneous neuronal activity and therefore can increase activity in a more physiological manner. tDCS also offers a valuable practical advantage as investigators and study subjects can be reliably blinded, therefore allowing for well-controlled experiments (Gandiga, Hummel, & Cohen, 2006).

Nitsche and colleagues showed that anodal tDCS of the human motor cortex elicits prolonged cortical excitability increases (Nitsche & Paulus, 2000, 2001), most likely by inducing sub-threshold neuronal membrane depolarization (Purpura & McMurtry, 1965). Moreover, it has been shown that tDCS evokes after-effects that are NMDA receptor dependent (Liebetanz, Nitsche, Tergau, & Paulus, 2002) and that share some similarity with the LTP and LTD presumed to underlie learning processes (Rioult-Pedotti et al., 1998). These studies suggest that anodal tDCS has the potential to improve rehabilitative learning by focally increasing cortical excitability and modulating NMDA receptor dependent plasticity in areas that facilitate motor recovery during behavioral training.

Indeed studies suggest that tDCS enhances motor learning in healthy subjects (Boggio et al., 2006) and stroke patients (Fregni & Pascual-Leone, 2005; Hummel & Cohen, 2005). These findings have been confirmed by others and extended to language function in normal subjects (Floel & Cohen, 2007) and in patients with aphasia (Monti et al., 2008). There are also preliminary reports in healthy subjects of improved verbal fluency (Iyer et al., 2005) and verbal working memory abilities (Fregni, 2005b) associated with combined tDCS/behavioral training as compared to behavioral training alone. Thus tDCS may serve as a simple yet powerful supplementary tool for enhancing learning associated with physical rehabilitation (Bolognini, Pascual-Leone, & Fregni, 2009).

Conclusions

Functional motor recovery following stroke remains limited, with many adult stroke survivors never approaching full functional recovery. Interestingly, children exhibit better rehabilitation outcomes when compared to adults suffering from comparable brain injuries, though the reasons why remain unclear. Here, we discuss possible biological explanations for the inadequate functional recovery commonly observed in adults as well as the remarkable recoveries commonly observed in children.

It is clear from evidence on the molecular, cellular, and systemic levels that neuroplasticity is not absent in adult patients following stroke, and, to the contrary, that a multitude of mechanisms geared toward promoting brain reorganization are enhanced following injury. It is therefore unlikely that an inherent lack of neuroplasticity alone could fully account for relatively poor functional recovery in adults. We suggest that decreased motor recovery in adults is in part due to inhibitory neuronal interactions, such as transcallosal inhibition, that serve to optimize motor performance as the brain matures. Following stroke, these inhibitory interactions pose rigid barriers to recovery by inhibiting activity in the affected regions and hindering recruitment of compensatory pathways. We suggest that children exhibit better rehabilitation outcomes in part because they have not fully developed the inhibitory interactions that obstruct functional recovery in adults.

We conclude with a discussion of how TMS and tDCS can be used to enhance patient care and motor recovery following stroke. TMS can be used as a diagnostic tool for recording neurophysiological measures and tracking improvement following stroke. Both techniques can be employed to effectively restore interhemispheric balance to the brain by inhibiting the inhibitory systems that develop as one ages, effectively creating brain states that might be partially equivalent to a “premature” state. Both techniques can also enhance synaptic plasticity and re-learning of motor skills following stroke. In fact we have recently compared intensive motor training with sham tDCS vs. intensive motor training with tDCS aimed to reduce transcallosal inhibition from the unaffected to the affected motor cortex. In this study we have observed that intensive motor training alone has inferior results as compared to the group that also received tDCS to
decrease transcallosal inhibition. Indeed the additional beneficial behavioral effects were correlated to a decrease in transcallosal inhibition (data not published).

Additional research is needed to support the conclusions we discuss here. While we present a large literature that demonstrates robust neuroplasticity in adults, more direct comparisons quantifying the extent of neuroplasticity in children compared to adults (on molecular, cellular, and systemic levels) would be needed to rule out the possibility that differences in neuroplasticity across age can account for rehabilitation differences. Furthermore, a properly controlled experiment in healthy subjects could be proposed to examine our hypothesis that poor functional recovery in adult stroke patients compared to pediatric stroke patients is largely due to maladaptive consequences of increased transcallosal inhibition in adults. Such an experiment might involve TMS induction of intracortical inhibition and facilitation in adults and kids as they learn a motor task, and measuring how transcallosal inhibition correlates with motor learning throughout the experiment. Future studies looking at the effects of rTMS and tDCS on motor recovery following stroke, should combine these noninvasive brain stimulation techniques with different rehabilitation approaches, employ varying parameters, and utilize greater numbers of subjects, so that we can reliably evaluate these techniques and generate optimal rehabilitative therapies.

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References

populations in the macaque monkey are specified by their unique cortical origins, spinal terminations, and connections. *Cerebral Cortex*, 4, 166-194.


