



Consensus: Can transcranial direct current stimulation and transcranial magnetic stimulation enhance motor learning and memory formation?

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Noninvasive brain stimulation has developed as a promising tool for cognitive neuroscientists. Transcranial magnetic (TMS) and direct current (tDCS) stimulation allow researchers to purposefully enhance or decrease excitability in focal areas of the brain. The purpose of this article is to review

Funding provided by the National Institutes of Health.

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Submitted June 5, 2008; revised July 29, 2008. Accepted for publication August 2, 2008.

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doi:10.1016/j.brs.2008.08.001

57 information on the use of TMS and tDCS as research tools to facilitate motor memory formation, motor
 58 performance, and motor learning in healthy volunteers. Studies implemented so far have mostly
 59 focused on the ability of TMS and tDCS to elicit relatively short-lasting motor improvements and the
 60 mechanisms underlying these changes have been only partially investigated. Despite limitations,
 61 including the scarcity of data, work that has been already accomplished raises the exciting hypothesis
 62 that currently available noninvasive transcranial stimulation techniques could modulate motor learning
 63 and memory formation in healthy humans and potentially in patients with neurologic and psychiatric
 64 disorders.

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66 **Q1 Keywords** ■

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 69 Within the past 2 decades noninvasive brain stimulation
 70 has been used as a probe to modulate attention, memory,
 71 motor, and language functions in humans.¹⁻⁹ TMS and tDCS
 72 can enhance or decrease excitability in target cortical regions
 73 depending on the parameters of stimulation used.¹⁰⁻¹³ TMS,
 74 and to a lesser extent tDCS (specific differences are
 75 described by Nitsche et al in this issue of *Brain Stimulation*)
 76 have been used as an interference technique (“virtual
 77 lesion”) for understanding brain-behavior interactions and
 78 to explore possible cause-effect links between altered activity
 79 in specific brain areas and particular behaviors.^{14,15}
 80 Improved understanding of the involvement of a brain
 81 region in a type of behavior was followed by attempts to
 82 modify activity in this area to secondarily influence perfor-
 83 mance, learning, and memory functions.²⁻⁹ In this chapter
 84 we summarize the results from studies that aimed at eliciting
 85 improvements in motor performance and motor learning
 86 in healthy humans.

88 **Motor learning**

91 Formation of motor memories is required for learning the
 92 motor skills in daily life.^{16,17} It is helpful to distinguish
 93 studies focused on the process of acquisition, consolidation,
 94 and long-term stability (also referred to as retention) of a
 95 new motor skill^{8,18,19} from those that evaluate the return
 96 to baseline levels of performance in response to external
 97 perturbations.²⁰⁻²³ An example of adaptation to an external
 98 perturbation is the response to directional errors in visually
 99 guided reaching movement caused by prism glasses:^{22,24}
 100 with practice, performance returns to the “baseline” level.
 101 Importantly, adaptation may not require the acquisition of
 102 new motor synergies or movement patterns, as it engages
 103 movements that were achieved throughout life.

104 In contrast to adaptation, acquisition of a new motor
 105 skill involves the acquisition of new movement qualities
 106 and/or muscle synergies that enhance performance beyond
 107 preexisting levels. Skills seem to take longer to acquire than
 108 adaptation and sometimes do not reach plateau levels after
 109 years (ie, learning to play piano or basketball).^{20,25-27} In
 110 engineering terms, adaptation may be modeled as error-
 111 based learning, whereas, for example, motor skill learning
 112 is better modeled in terms of reward-based signals.^{28,29}

113 However, it should be kept in mind that an overlap of
 114 reward- and error-based learning is possible.

115 Consolidation refers either to stabilization (reduced
 116 susceptibility to retrograde interference) or offline improve-
 117 ments.³⁰⁻³³ Consolidation processes can depend on the type
 118 of task, the time between the end of practice and the testing
 119 of recall, and the presence or absence of sleep. Offline im-
 120 provements, for instance, in the ability to perform a finger
 121 opposition task, correlated with the amount of time spent in
 122 REM sleep.³⁴ Similarly, offline improvements in a motor
 123 sequence learning task are sleep dependent when individ-
 124 uals are aware of the underlying sequence.³⁵ However,
 125 when individuals have little awareness for the sequence,
 126 offline improvements are able to develop over waking or
 127 over a night sleep.³⁵ Potentially, the effect of individuals’
 128 awareness on offline learning is mediated by their declara-
 129 tive knowledge for the sequence: disrupting declarative
 130 knowledge for the sequence can induce improvements
 131 over wake.³⁶ In adaptation studies, the successful return
 132 to baseline performance after the perturbation occurs often
 133 within one session, and therefore the possibility of offline
 134 improvements across days has not been thoroughly tested,
 135 although savings, an increase in the rate of readaptation,
 136 could be considered a form of offline learning.³⁷ One
 137 exception to this general statement is the study of Huber
 138 et al³⁸ in which overnight improvements in performance
 139 of a motor adaptation paradigm in which subjects had to
 140 adapt to a visual perturbation of a reaching movement cor-
 141 related with increased less than 4 Hz activity during slow
 142 wave sleep. This activity is thought to reflect oscillatory
 143 changes in neuronal membrane potentials.³⁸

144 As stated previously, another form of consolidation is
 145 stabilization, that is, maintenance of practice-induced per-
 146 formance improvements or skill (in opposition to forgetting
 147 or to offline improvements). After the end of a practice
 148 period, procedural memories for a task A may display
 149 different degrees of strength to interference. A classical
 150 approach to evaluate this strength is to introduce a task B as
 151 a source of interference and subsequently test the subject’s
 152 ability to perform task A.^{37,39,40} Interference to recall a
 153 newly learned motor task A by practicing a different motor
 154 task B has been described as “retroactive interference.”
 155 Retroactive interference has a well-described time course
 156 diminishing with the length of the time interval between

156 the end of practice of task A and the application of the
157 interfering task B, becoming virtually absent after
158 6 hours.^{37,40,41} Stabilization over hours after learning
159 dynamic adaptation tasks has been well-documented.^{41,42}
160 Of note, Goedert and Willingham⁴⁰ showed that for motor
161 sequence learning offline stabilization does not occur.
162 Whether 15 minutes or 24 hours passes in between task A
163 and B, task B continues to exert retrograde interference.⁴⁰
164 However, this finding might be task specific.

165 It is important to keep in mind that during motor
166 learning, both kinds of consolidation, offline learning and
167 stabilization, are likely occurring.^{43,44} As stated previously,
168 it is unclear if offline enhancement occurs for adaptation
169 tasks as consolidation studies of adaptation have focused
170 on stabilization. Other factors that may influence our ability
171 to assess the stability of a procedural memory include the
172 end point measure used (speed, accuracy, or speed accuracy
173 ratios related to a motor task) and the practice sched-
174 ule:^{45,46} for instance, when a skill is acquired through inter-
175 leaved rather than blocked practice schedules, motor
176 memories may become more resistant to interference.

177 One important theoretical point to highlight is the differ-
178 ence between measurements of motor performance and
179 motor skill. Improvements in speed or in accuracy of
180 performance of a motor action have been often reported in
181 isolation in the literature, occasionally indicating that
182 changes in one of these two measures occurred in the absence
183 of changes in the other. Such changes have been reported as
184 changes in skill. It would be important to keep in mind that
185 motor skill cannot always be reliably surmised from changes
186 in only one of these two measures. Skill may be better
187 described as a change in the speed-accuracy trade-off, which
188 is task dependent. Taking into consideration this issue would
189 help future investigators avoid concluding a change in skill
190 (skill improvements) when in fact subjects have only moved
191 along the same speed/accuracy trade-off curve.

194 Noninvasive brain stimulation

196 Noninvasive brain stimulation has been used to identify the
197 functional relevance of particular brain regions in motor
198 learning and facilitate activity in specific cortical areas
199 involved in motor learning in an attempt to improve motor
200 function.

203 Functional role of the primary motor cortex 204 in motor learning as studied with 205 noninvasive cortical stimulation

208 Motor learning is associated with functional changes in a
209 distributed network that includes the primary motor,
210 premotor and supplementary motor cortices, the cerebel-
211 lum, thalamic nuclei, and the striatum.^{18,47-50} Most TMS

212 and tDCS studies performed so far to study the role of
213 motor areas in motor learning have focused on M1.

214 *Role of M1 in encoding of an elementary motor memory:*
215 Butefisch et al⁵¹ showed that the synchronous application
216 of single-pulse TMS to M1 contralateral to a hand practic-
217 ing a thumb abduction task enhanced the ability of healthy
218 subjects to encode an elementary and short-lasting motor
219 memory in the primary motor cortex.² Importantly, this
220 effect was evident when M1 was stimulated in synchrony
221 with the training motions but not when applied in between
222 training movements. A second important finding from this
223 study was that synchronous stimulation of the "resting"
224 M1 with the training motions in the ipsilateral hand,
225 cancelled training effects on motor memory formation,
226 consistent with the hypothesis that interhemispheric inter-
227 actions between M1s contribute to motor memory forma-
228 tion.² It has been proposed that formation of motor
229 memories within M1 could represent a first step in the
230 more complex chain of events leading to improve a motor
231 skill, but it should not be interpreted as motor learning
232 *per se* as skill improvements above naïve levels are typi-
233 cally not seen with this particular paradigm of motor
234 memory formation.

235 *Role of M1 in motor adaptation:* As discussed previ-
236 ously, the concept of motor adaptation refers to learning
237 to adjust to external perturbations.^{21-23,52} In these experi-
238 ments, subjects adjust their motor behavior to compensate
239 for a particular perturbation to maintain a stable perfor-
240 mance.²¹ TMS and tDCS have been used to evaluate the
241 role of M1 in motor adaptation. In general, stimulation
242 over M1, using parameters that decrease excitability in
243 that region like 1-Hz TMS, have been applied before or
244 during adaptation paradigms (to evaluate its functional
245 relevance for encoding of the necessary adjustments to
246 compensate for the perturbation).^{23,53} In one study, single
247 TMS pulses applied to M1 at 120% of resting motor thresh-
248 old (RMT) of the first dorsal interosseus muscle immedi-
249 ately after the end of each trial while adapting to a
250 perturbation in the form of a visuomotor rotation did not
251 impact adaptation, but caused faster deadaptation (forget-
252 ting) within the same session relative to single pulses ap-
253 plied 700 milliseconds after the end of each trial or
254 relative to PMd stimulation.⁵³ In another study, 1-Hz
255 rTMS applied to M1 at 90% of biceps RMT before force
256 field adaptation did not affect the participants' adaptation
257 *per se*, but impaired retention relative to control subjects
258 (who did not receive any rTMS) as tested the following
259 day when subjects were exposed to the same force field
260 to which they had previously adapted.⁵⁴ On the other
261 hand, Baraduc et al²³ did not find a deleterious effect of
262 1-Hz TMS applied over M1 on adaptation to a dynamic
263 force field. Potential areas of interest that remain to be
264 investigated in more detail include the role of motor areas
265 other than M1 in motor adaptation.

266 *Role of M1 in motor skill learning:* As discussed previ-
267 ously, motor learning may (and often does) continue after

the end of practice periods, referred to previously as consolidation in the form of offline learning. The role of M1 has been investigated in the process of acquisition and consolidation of motor skills.^{13,50,55} Muellbacher et al¹³ reported in an influential study, that 1-Hz rTMS over M1 at 115% of flexor pollicis brevis RMT applied immediately before a single-session practice of a thumb-to-finger opposition task did not disrupt within session improvements in speed and muscle force generation but had deleterious effects on retention of these improvements as tested the following day relative to stimulation applied 6 hours after practice or when applied to other cortical areas such as the occipital cortex or the left DLPFC. These results were interpreted as supportive of the view that M1 plays a functionally relevant role in consolidation of explicit motor memories.¹³ The role of M1 was also explored in motor sequence learning⁵⁶ by using a modified version of the serial reaction time task (SRTT).⁵⁷ 1-Hz rTMS was applied over M1 immediately after training when subjects practiced the task early in the morning or late in the evening (different groups). The end point measure was offline enhancement in performance of the task 12 hours later (the evening of the practice day with no sleep in between in the first group and the following morning after a normal night sleep in the second group). It was reported that offline enhancements of the learned task were disrupted in the first group (no sleep) but not in the second group. The authors interpreted the result as indicative of different consolidation processes depending on how close sleep is to the practice period.⁵⁶ For a more detailed review of the effects of virtual lesion studies on motor learning, please refer to¹⁵. On the other hand, rTMS and tDCS have been used in an attempt to facilitate motor learning, the focus of this review.

Attempts to facilitate motor learning by noninvasive cortical stimulation

On the basis of human neuroimaging studies, it was proposed that application of noninvasive stimulation with parameters that enhance motor cortical excitability could secondarily facilitate motor learning. One key structure in the distributed network engaged in motor sequence learning is the primary motor cortex (M1).^{13,58} Within M1, the extent of cortical reorganization associated with motor training correlates with performance improvements.⁵⁹ The interaction between the two M1s appears to play an important role in motor control in general,^{20,60-64} and in motor sequence learning in particular.^{64,65} However, the specific way in which these interactions operate during motor learning remain to be determined. According to these interactions, it would be theoretically possible to facilitate motor learning processes in which M1 is involved by enhancing excitability in the "learning" M1⁵ or by decreasing excitability in the "resting" M1.⁶⁶⁻⁶⁸ The intrinsic intracortical

mechanisms by which these oversimplified models may operate remain to be identified^{64,69,70} (see for discussion chapters by Walsh et al., Di Lazzaro et al., Berardelli et al.).

Several investigators proposed that noninvasive cortical stimulation that enhances excitability in the M1 contralateral to a training hand^{5,71} might result in varying degrees of improvement in motor function in healthy humans. Kim et al⁷² reported that 10-Hz rTMS at an intensity of 80% of RMT applied over M1 during practice of contralateral sequential finger movements resulted in improved motor sequence learning (as measured by target accuracy and speed on a sequential key press task) relative to sham stimulation. Anodal tDCS applied over M1 during practice also led to improvements in: (1) the number of correct key presses in a sequential finger movement task in a polarity-specific manner since cathodal tDCS failed to induce this effect⁷³; (2) performance of a visuomotor coordination task that were transient (~5 minutes)⁷⁴; (3) reaction times in the sequence blocks relative to the random blocks in the SRTT,⁵⁷ in which subjects learn a sequence of 12 key presses without explicit awareness⁷⁵; and (4) performance of the Jebsen Taylor Hand function test (JTT), a task often used in stroke research that mimics activities of daily living such as lifting cans and picking up small objects.⁷⁶ Interestingly, this effect has been reported as present in the nondominant hand only in young healthy adults.⁷⁷ Of note, tDCS in these two articles was applied after subjects reached stable JTT performance, likely reflecting a tDCS-induced performance improvement beyond a plateau level. It is not known if application of noninvasive cortical stimulation during the learning period of the task (before it reaches an asymptote) could speed up or enhance learning of the task.

In contrast to studies that focused on application of TMS or tDCS to the M1 contralateral to a practicing hand, the application of 1-Hz rTMS to the M1 ipsilateral to a training hand results in: (1) increases in motor cortical excitability of the opposite M1⁶⁶⁻⁶⁸ relative to sham stimulation, and (2) improvements in motor sequence learning⁶⁵ relative to stimulation of the contralateral M1, ipsilateral premotor area, or vertex (Cz). One important consideration is that the effects of stimulating M1 with either TMS or tDCS are likely to be dependent on the complexity of the task. For example, performance of relatively simple repetitive finger abduction movements was not improved by high-frequency rTMS over the "learning" M1,⁷⁹ whereas more complex sequential motor tasks or encoding of a motor memory did improve (discussed previously). Similarly, another study showed no performance improvement in a task engaging single finger tracking motions when the ipsilateral M1 was stimulated with 1-Hz rTMS during practice.⁸⁰

In contrast to studies focusing on motor sequence learning or motor performance, we are not aware of studies that used TMS or tDCS in an attempt to facilitate motor adaptation.

Caveats and future directions

Although the previous paragraphs depict a relatively consistent and homogenous picture on the effects of up- and down-regulation of excitability within M1 on motor learning and motor memory formation, several caveats should be kept in mind. First, induction of a “virtual lesion” or enhancement of activity in one cortical area may result in behavioral changes through specific effects on that area or secondarily through distant effects on other interconnected cortical areas. Second, the discussion of results in this article (as well as those of specific physiologic interactions across cortical regions⁶¹) assume, in general, that the surface of the brain is a smooth sphere, often neglecting that the folding of the cortex may result in hyperpolarization of neurons on one side of a gyrus but depolarization on the other. Third, the history of activity in the stimulated cortex may be of considerable importance. For example, the effects of stimulation may differ substantially if applied to a “fatigued” or to a “well-rested” cortex. In one example, 1-Hz rTMS may induce facilitatory effects if acting on a cortex that has been previously inhibited by cathodal tDCS.^{81,82} This phenomenon, referred to as homeostatic plasticity or metaplasticity and discussed elsewhere,⁸³ may potentially impact motor learning.^{84,85} Fourth, although most of the work in the field focused on studying the effects of stimulation over M1, some reports indicated that stimulation of the dorsal premotor or the lateral or medial prefrontal cortex failed to induce overt reaction time improvements in the SRTT task.⁷⁵ Similarly, anodal tDCS applied over the primary visual cortex (V1) did not improve performance of a visuomotor tracking task in healthy subjects.⁷⁴ However, findings indicating a lack of effect of stimulation in a particular site are not proof that the area is not involved. The most parsimonious interpretation of these findings is that more elaborated, hypothesis-driven behavioral paradigms or stimulation strategies may be necessary to study the functional role of these cortical regions in motor learning and memory formation. Alternatively, it is possible that the “threshold” for facilitating motor learning by M1 stimulation is lower than by stimulation of other cortical areas, an issue to be investigated in future experiments. Fifth, one caveat of many previous investigations has been the focus on short-term improvements in performance. More experiments are required to assess the effects of repeated applications of TMS or tDCS in association with multiple training sessions, their interaction with specific motor learning stages and tasks, and the extent to which these performance improvements are retained in the long term. Finally, it should be kept in mind that the effects of TMS and tDCS cannot be assumed to be the same. The mechanisms underlying the effects of each technique on motor cortical function are very likely to differ, as discussed by Nitsche et al in this issue of *Brain Stimulation*, and so will the effects on motor cortical networks and

behavioral consequences of its application. Clearly, more work is needed in this area.

Summary

In summary, the scarce studies performed so far point to the encouraging conclusion that noninvasive brain stimulation can contribute to the understanding of mechanisms underlying motor learning and motor memory formation and raise the exciting hypothesis that this increased understanding could in the future result in the development of new strategies to enhance specific stages of learning and memory processing in healthy humans and in patients with brain lesions, as discussed by Gerloff et al in this issue of *Brain Stimulation*.

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