Exercise-mediated locomotor recovery and lower-limb neuroplasticity after stroke

Larry W. Forrester, PhD;1–2* Lewis A. Wheaton, PhD;3 Andreas R. Luft, MD4

1Department of Veterans Affairs (VA) Maryland Health Care System, Research Service, Baltimore, MD; 2Department of Physical Therapy and Rehabilitation Science, University of Maryland School of Medicine, Baltimore, MD; 3VA Maryland Health Care System, Baltimore, MD; 4Hertie Brain Institute for Clinical Brain Research, University of Tübingen, Tübingen, Germany

Abstract—Assumptions that motor recovery plateaus within months after stroke are being challenged by advances in novel motor-learning-based rehabilitation therapies. The use of lower-limb treadmill (TM) exercise has been effective in improving hemiparetic gait function. In this review, we provide a rationale for treadmill exercise as stimulus for locomotor relearning after stroke. Recent studies using neuroimaging and neurophysiological measures demonstrate central nervous system (CNS) influences on lower-limb motor control and gait. As with studies of upper limbs, evidence shows that rapid transient CNS plasticity can be elicited in the lower limb. Such effects observed after short-term paretic leg exercises suggest potential mechanisms for motor learning with TM exercise. Initial intervention studies provide evidence that long-term TM exercise can mediate CNS plasticity, which is associated with improved gait function. Critical needs are to determine the optimal timing and intensities of TM therapy to maximize plasticity and learning effects.

Key words: gait, gait training, hemiparesis, locomotor, lower limb, motor control, motor learning, neuroplasticity, neurorehabilitation, rehabilitation, stroke, treadmill exercise.

INTRODUCTION

Approximately 700,000 strokes occur annually in the United States; 50 percent of the 550,000 survivors experience residual hemiparesis and approximately 165,000 of those individuals have mobility deficits requiring assistance with walking [1–3]. In this population, hemiparetic gait is a major problem that limits mobility, increases risk of falls, and imposes higher energy demands for basic daily activities [4–5]. Gait deviations due to hemiparesis are well documented, in terms of both clinical manifestation and biomechanical analyses [6–7]. Classic models of stroke recovery indicate that improvements in both upper- and lower-limb motor function plateau between 3 and 6 months poststroke [8]. Recent studies have challenged this assumption by suggesting that specific training interventions that target use of the hemiparetic limbs can improve motor control and neural plasticity. The research community now

Abbreviations: CNS = central nervous system, EMG = electromyography, FES = functional electrical stimulation, fMRI = functional magnetic resonance imaging, M1 = primary motor cortex, MEP = motor-evoked potential, MRCP = movement-related cortical potentials, NIRS = near-infrared spectroscopy, PAS = paired associative stimulation, PBWS = partial body-weight suspension, RR&D = Rehabilitation Research and Development, S1 = primary somatosensory cortex, SCI = spinal cord injury, SMA = supplementary motor area, SMC = sensory motor cortex, TA = tibialis anterior, T-AEX = treadmill aerobic exercise, TM = treadmill, TMS = transcranial magnetic stimulation, VA = Department of Veterans Affairs.

*Address all correspondence to Larry W. Forrester, PhD; University of Maryland School of Medicine, 100 Penn Street, Suite 115, Baltimore, MD 21201-1082; 410-706-5212; fax: 410-706-6387. Email: lforrester@som.umaryland.edu

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widely accepts that the central nervous system (CNS) comprises inherently plastic neural networks that are continuously amenable to reorganization in the service of functional behaviors [9]. As a consequence, new therapeutic approaches seek to exploit experience-based CNS plasticity to mediate functional improvements. A common thread among most of these interventions is an adherence to the principles of motor learning, as defined by incorporating high volumes of task-oriented practice along with the added dimensions of goal setting and performance feedback [10].

Studies of therapies that improve function and induce neuroplasticity in hemiparetic upper limbs in human survivors of stroke have supported an emerging focus on developing new learning-based strategies for improving gait and balance function in individuals with lower-limb hemiparesis after stroke [11–17]. Here we review evidence that one particular mode of exercise, treadmill (TM) training as applied in a number of approaches, can be employed to improve gait function in survivors of stroke with residual hemiparesis. We will suggest that basic motor learning strategies can alter underlying neural mechanisms to improve hemiparetic function of the lower limb and may also be effective in recovery of walking ability after stroke. Following a brief overview of the rationale and early results from studies using TM training with stroke, we provide examples that illustrate the role of the CNS in lower-limb motor control and gait. Our focus then shifts to an overview of how the neurophysiology of lower-limb motor control is sensitive to short-term adaptations and rapid plasticity. Finally, we review the early evidence of central neuroplasticity underlying lower-limb function and gait using long-term TM training protocols.

RATIONALE FOR TREADMILL LOCOMOTOR LEARNING AFTER STROKE

Findings from spinalized animal models demonstrate that walking without supraspinal inputs can occur when the animal is placed on a moving TM [18]. Thus, several investigations have studied TM training as a means to improve locomotor function in subjects with incomplete spinal cord injury (SCI) and stroke. Visintin et al. first adapted the findings from spinalized animals to human experiments, reasoning that activation of subcortical neural structures by TM walking could provide a physiological stimulus for recovery of gait function [19–20]. These studies support the rationale that TM-generated stepping patterns in neurologically injured humans can help deliver repetitive sensory inputs to the spinal cord, which in turn could mediate locomotor learning and neural plasticity through a process of sensory motor integration [21]. Additional feasibility for this idea was shown in a study of the immediate effects of the TM stimulus on hemiparetic gait patterns in naive subjects with chronic stroke [22]. While controlling for walking speed, paretic limb stance-swing parameters and loading impulse immediately became more symmetrical on the TM compared with usual overground walking. Analyses of electromyography (EMG) activation patterns showed that this symmetry was not an artifact of TM-induced mechanical perturbations, as timing of EMG bursts shifted significantly within the paretic gait cycle [23]. Thus, untrained individuals with hemiparesis can alter how they walk during a brief exposure to the TM stimulus. A question then arises as to whether an adequate amount of practice would promote lasting changes in their gait function. If so, is the effect reflected in measures of lower-limb motor control and central neural plasticity?

TREADMILL-BASED EXERCISE TRAINING IMPROVES GAIT FUNCTION

The initial studies with human SCI and subacute stroke subjects used TM training in conjunction with partial body-weight suspension (PBWS). In a randomized study of more severely impaired subjects with subacute stroke, Barbeau and Visintin found TM with PBWS to be more effective than TM without PBWS for improving selected mobility outcomes in those subjects with more severe motor deficits (i.e., <0.2 m/s walking velocity and Berg Balance scores <15) [24]. By week 6 of training, 79 percent of subjects were able to train at 0 percent PBWS. In a noncontrolled 3-week study, 25 PBWS TM training sessions improved mobility scores and gait temporal-distance parameters in nine nonambulatory stroke subjects [25]. PBWS was not required after day 6 of training in seven of these nine cases. Similar results were reported in a follow-up study using the same approach in an A-B-A design [26]. These studies indicate an important role for PBWS as a bridge to full-weight-bearing TM exercise, particularly in subjects more severely affected.
Other therapeutic approaches have been adapted from the original PBWS TM studies to include novel applications of robotically facilitated gait training with and without the TM and also with augmentation by functional electrical stimulation (FES). These approaches emphasize new gait therapies for nonambulatory patients with severe paresis after stroke or SCI. The Lokomat (Hocoma AG; Volketswil, Switzerland) is a robotic gait trainer that integrates PBWS and TM with actuated hip and knee orthoses to emulate normal walking patterns [27]. A recent randomized crossover study found that subjects with hemiparesis made greater improvements in gait function and lower-limb impairment measures following periods of Lokomat training compared with equal periods of conventional physical therapy [28]. Husemann et al. showed that 4 weeks of Lokomat and usual therapy improved the functional ambulation category for subjects in the acute phase of stroke as well as in those who received equal amounts of usual therapy only. However, the Lokomat group increased paretic single support times in overground walking, gained more muscle mass, and lost more fat compared with the controls, who gained fat mass [29]. Hesse et al. developed and tested an electromechanical gait trainer to move the legs in a manner physiologically similar to walking [25]. A study of survivors of stroke in the subacute phase of recovery showed that the electromechanical gait trainer was as effective as therapist-assisted PWBS TM training for improving gait function [26].

The gait trainer has also been used in conjunction with FES applied to knee extensors and ankle dorsiflexors in nonambulatory subjects with hemiparesis, for comparing the possible benefits of combined treatment versus either usual therapy or gait trainer alone in a 4-week intervention [30]. The gait trainer with FES group and gait trainer only group improved more than controls, but the two gait trainer methods did not differ. In another A-B-A design study of a 9-week protocol, FES was used to augment PBWS TM therapy in a small sample of subjects with chronic stroke. Gait speed, cadence, and stride length increased significantly after the introduction of FES, and gait speed declined when FES was discontinued during a final phase of PBWS TM training only [31]. The potential for early intervention to enhance gait function by combining TM training with robotic assistance and/or FES is promising; however, further studies are needed to delineate optimal methods.

The question of how to optimize TM training for improving gait function after stroke is important but unsettled. We have investigated yet another approach to TM training during the chronic phase after stroke by emphasizing progressive cardiovascular demands over a 6-month program. Improved floor walking speeds, economy of gait, and cardiovascular fitness were reported for subjects with chronic hemiparesis after 6 months of TM aerobic exercise (T-AEX) training using handrail support without PBWS (Figures 1–2) [32]. A reference control group spent equivalent time performing stretching exercises. An important feature of the T-AEX protocol was the emphasis on aerobic conditioning, with increased TM walking duration and velocity to maintain 60 percent of heart rate reserve, a key indicator of exercise intensity [33–34]. While the primary focus was on cardiovascular conditioning, improvements were also found in fundamental gait parameters, indicating that T-AEX can differentially affect step lengths and walking cadence to achieve increased velocity [35]. As well, the double stance times

![Figure 1. Mean percent change in 6-minute walk distance in treadmill aerobic exercise (T-AEX) group (solid line) and reference control (R-CONTROL) stretching groups (dashed line). Significant group-by-time interaction occurred in 6 min walk distance by repeated measures analysis of variance (*p < 0.02) with progressive gains across 6-month intervention period (*p < 0.05). Values are mean ± standard error. Source: Reprinted by permission from Macko RF, Ivey FM, Forrester LW, Hanley D, Sorkin JD, Katzel LI, Silver KH, Goldberg AP. Treadmill exercise rehabilitation improves ambulatory function and cardiovascular fitness in patients with chronic stroke: A randomized, controlled trial. Stroke. 2005;36(10): 2206–11. [PMID: 16151035]]
decreased, suggesting improved postural stability during weight shifts between the legs. Consideration of these changes in gait parameters and cardiovascular fitness together highlights the potential for T-AEX to translate improved gait function into capacities needed for sustained mobility in daily living and may help define clinically significant outcomes.

Other investigators have begun to focus on training intensity, which may involve manipulation of the velocity, duration, grade of incline, and concentration (massing) of practice. Sullivan et al. reported that after a 4-week PBWS training program, survivors of stroke who trained at faster TM velocities had greater increases in the criterion test of self-selected floor walking velocity \( [36] \). In a randomized controlled trial, Pohl et al. systematically applied higher velocities to elicit greater improvements in overground walking parameters (velocity, cadence, stride length, functional ambulation category) after 4 weeks of TM training, compared with two reference groups receiving 20 percent speed increases or no speed increases \([37]\). Others are now looking at the use of TM grade to intensify the training stimulus, with one report showing increases in heart rate and improved gait pattern symmetry and longer stride lengths when grade is increased to 8 percent for subjects with hemiparesis \([38]\). In yet another variation of TM training, a split-belt TM altered hemiparetic gait patterning through differential belt speeds \([39]\). Adaptations as after-effects in gait kinematics were observed in subjects with stroke, and these persisted briefly when subjects were transitioned immediately to overground walking. Although not yet proven durable, this observation suggests that the mechanical TM stimulus is affecting CNS motor planning of gait.

Few investigations have directly linked TM training to overground walking. Plummer et al. have been proponents of coupling PBWS TM training with transfer to the specific task of overground walking, immediately reinforced by cueing appropriate arm and stepping actions \([40]\). Their approach is grounded in neurally based functional requirements for walking, and their pilot data suggest that such an approach is safe and feasible for improving gait function among individuals who are moderately to severely impaired. We also have begun to look at the question of carryover from TM training to independent walking and report preliminary data on gait pattern changes after 6 months of T-AEX \([35]\). A key finding was that velocity improvements in unassisted 8-meter walks were due to a combination of increased stride length and frequency. Importantly, the training did not alter interlimb symmetry in either step times or step lengths; hence, both limbs appeared to amplify the preexisting hemiparetic pattern to improve overall gait function.

In the context of defining optimal training approaches, little is known about the interactions between deficit severity and any of these various training parameters. Individuals with stroke tend to have multiple comorbid conditions that can affect participation in TM training. This issue is now receiving closer attention. For example, in their pilot feasibility study, Plummer et al. stratified subjects with stroke according to self-selected walking velocity (<0.4 m/s vs. >0.4 and <0.8 m/s) \([40]\). The subjects who were moderately impaired made clinically meaningful gains after 24 sessions and the subjects who were severely impaired were improving, but not to a clinically meaningful level after the full 36-session program. This finding...
starts to provide a basis for constructing individualized therapy regimens based on ambulatory function.

Taken together, these studies indicate that concentrated practice through TM exercise training can improve gait function in survivors of subacute and chronic stroke. Mechanistically, they suggest that repetition of an effective gait pattern/rhythm may be critical to restoring gait function. However, also likely is that the long-term TM exercise affects a number of other processes besides learning a more functional gait pattern, including biological responses in peripheral muscle, balance control, and self-efficacy related to fall risk. Thus, a complete understanding of what transpires during any of these TM training regimens is very difficult to realize as we consider the potential mechanisms for improving hemiparetic gait. In the following sections, we focus on the emerging evidence that, like the upper limb, central neural plasticity is a likely mechanism underlying lower-limb functional recovery after stroke and that TM training can be a viable motor-learning stimulus for triggering that response.

CNS ROLE IN LOWER-LIMB MOTOR CONTROL AND GAIT

The neurophysiology of lower-limb motor control and its impact on locomotor recovery has become another focus for poststroke rehabilitation. Corticospinal connectivity to lower-limb musculature that determines ambulatory performance capacity is crucial to locomotor efficiency and recovery of basic activities of daily living. Studies of gait recovery after incomplete SCI and during normal motor development strongly suggest that improvement of human walking depends on enhanced cortical input [41]. In this section we summarize recent findings that employ a number of methods used in neurophysiological studies of upper-limb motor control to explore the central neural mechanisms of lower-limb motor control.

One noninvasive method to investigate lower-limb neurophysiology is transcranial magnetic stimulation (TMS), in which motor-evoked potentials (MEPs) are evaluated in the lower-limb musculature for characterizing aspects of the corticospinal connectivity that may underlie control of gait. Prolonged MEP latencies indicate descending pathway injury [42–43]. In the subacute phase of stroke, the ability to elicit lower-limb MEPs predicts improved long-term hemiparetic leg recovery [44–45]. Like several studies that show significantly reduced MEP responses in the paretic arm or hand compared with responses in the nonparetic limb [46–49], TMS reveals decreased excitability to the paretic leg relative to the nonparetic leg [50]. These effects are noted mainly as increased motor thresholds, longer latencies, and reduced MEP amplitudes to paretic versus nonparetic quadriceps muscles. Furthermore, this effect was observed in individuals with a variety of lesion locations, illustrating the fundamental impairment of corticospinal connectivity associated with residual lower-limb weakness and hemiparetic gait.

A number of investigations with nondisabled individuals have used TMS to demonstrate the role of corticospinal connectivity in the control of walking. Using a specialized mounting apparatus to fix coil position, Schubert et al. applied TMS stimulations to the cortex during TM walking, showing that corticospinal excitability to ankle musculature was differentially affected by the phase of the gait cycle [51]. Additionally, excitability effects were substantially greater on dorsiflexors as compared with plantar flexors. Capaday et al. used a similar approach to administering TMS during TM walking and reinforced these findings, highlighting the importance of corticospinal connections to the tibialis anterior (TA) during swing phase, compared with relatively reduced MEP responses in the soleus during stance [52]. Several other studies from Bo Nielsen’s group have elaborated on corticospinal contributions to gait [53]. Again, during active walking, TMS effects on H reflexes during the stance phase of the gait cycle were monitored to show that walking increases corticospinal excitability to ankle muscles, as evidenced by increases in H reflexes during walking but not under a controlled standing condition [54]. Furthermore, submotor threshold TMS delivered during walking caused suppression of the rectified EMG bursts from the TA and soleus muscles were suppressed, indicating that intracortical inhibitory responses were directly affecting the motor controlled of gait [55]. This protocol was modified to also show that long-latency stretch reflexes of the TA in nondisabled humans are at least partially modulated by transcortical circuits [56].

At least two studies have investigated brain activity during actual walking in nondisabled subjects. Fukuyama et al. used single photon emission computed tomography to show that several brain areas were active during over-ground walking in healthy subjects, including supplementary motor area (SMA), medial primary somatosensory cortex (S1), striatum, cerebellum, and visual cortex [57]. Activity across these distributed sites suggested that the brain is required to organize a complex flow of ongoing
sensory and motor information during normal independent walking. Miyai et al. used near-infrared spectroscopy (NIRS) to show that walking and foot flexion cause bilateral primary motor cortex (M1) and SMA activation, compared with contralateral M1 foci during isolated arm movements [58]. Miyai et al. also extended this method to a small cohort of nonambulatory subjects with hemiplegia to characterize cortical responses during PBWS TM walking [59]. They employed two different modes of therapist assistance: one assisted the swing of the paretic leg directly and the other used pelvic maneuvers to facilitate paretic swing indirectly. In both modes, the NIRS maps indicated activation in the medial primary sensory motor cortices (SMCs), with more activity seen in the nonlesioned hemisphere. Enhanced activation of the premotor and presupplementary motor areas of the lesioned hemisphere were also observed during gait. The pelvic assistance method produced generally greater cortical activations compared with directly moving the paretic swing leg. While having the limitation of a small sample size, this study demonstrates the feasibility of using therapist-augmented PBWS TM exercise to engage cortical networks. The study suggests that different therapeutic strategies may have distinct effects on the CNS.

Another area of focus is determining the effects of differing sensory modalities on CNS activity. This relates to the role of feedback as a requirement for motor learning, and whether certain types and quantities of afferent information enhance or impede the learning process and neuroplasticity. In a manner similar to that for the upper limb [60], the cortical processing for lower-limb motor planning in nondisabled subjects adapts to increased sensory inputs by increasing recruitment of parietal, motor, and premotor areas [61]. Greater sensory demands from combined visual and proprioceptive modalities evoked increased movement-related cortical potentials (MRCPs) during performance of a knee extension task, compared with single modalities and unconstrained knee movements, which evoked the least activity (Figure 3). This increase in MRCP is encouraging because nonprimary motor areas are known to be involved in stroke motor recovery [17]. The increase suggests that rehabilitation strategies that use an enhanced sensory environment may induce greater activation along the neuraxis to mediate improved lower-limb function.

More broadly, these methods for instantiating the role of central neural processing in regulating motor activity related to normal lower-limb function also provide a basis for assessing how the CNS may adapt to exercise-mediated training in individuals with stroke. In the next section we examine findings of adaptations in CNS activity due to short-term exercise exposures.

**RAPID-TRANSIENT PLASTICITY IN LOWER LIMB**

Beyond investigating the nature of CNS activity in control of lower-limb muscles and gait function, noninvasive techniques have also revealed aspects of rapid CNS plasticity after brief exposures to motor practice. To a limited degree, these efforts parallel upper-limb studies that demonstrate the potential for rapid changes in CNS excitability and task-specific cortical activation in nondisabled and stroke populations. A seminal study by Classen et al. reported rapid plasticity in control of the thumb muscles in nondisabled subjects, as TMS to the same location caused the CNS to encode the opposite kinematic response after as little as 20 minutes of repetitive thumb exercises in the opposite direction [62].

Corticospinal responses to different modes of short-term ankle exercise have been investigated in nondisabled subjects [63]. Recruitment curves from single-pulse TMS indicated that corticospinal excitability of the TA muscle increased after skill-based ankle training consisting of 32 minutes of volitional dorsif- and plantar flexion movements to track a target on a computer screen. Reference conditions with equal amounts of passive ankle movements or nonskilled volitional ankle movements did not show increased excitability. Another outcome was a decrease in intracortical inhibitory responses measured by paired-pulse TMS. Intracortical facilitation was not affected by the exercise. These results, along with no change on motor threshold levels and a negative finding in recruitment curves measured using transcranial electrical stimulation, were interpreted to suggest that the excitability changes due to skill-based exercise occurred at the cortical level.

Paired associative stimulation (PAS) has been used to investigate bidirectional corticospinal excitability of the hand muscles [64] by applying peripheral nerve stimulation to activate sensorimotor cortex within specified time windows around a pulse of TMS. When the afferent signals arrive at the cortex slightly ahead of the TMS impulse, excitability of the efferent pathways is enhanced. A recent study with nondisabled subjects examined the effects of PAS on TA responses during and following a 20-minute bout of TM walking at a moderate velocity (1.1 m/s) [65].
When peroneal nerve stimulation was timed to reach SMC approximately 5 ms before TMS and during the swing phase of ongoing TM walking, the posttest MEPs at the TA were significantly enhanced. When the TMS was administered before arrival of the afferent volley during walking, the posttest MEP amplitudes decreased compared with baseline. These results provide further evidence that sensory activation plays a key role in mediating CNS plasticity, which may be useful in rehabilitation of lower-limb function. One other small pilot study has shown potential for using the PAS approach in a therapeutic context for individuals with chronic stroke [66]. Although gait training exercise was not part of the 4-week intervention, PAS was applied 30 minutes a day for a total of 20 sessions. While the small subject sample showed mixed results on neurophysiological measures after the treatments, most subjects showed increased MEP amplitudes. Also, some participants improved in walking cadence and stride length. This improvement could indicate that PAS may augment experience-based plasticity mechanisms that mediate functional gains after task-oriented training. However, further investigations are needed to assess these potentials, as Uy et al. emphasize that only some of the functional and neurophysiological measures produced...
significant changes, likely because of the small sample size and differences in lesion characteristics [66].

The effects of TM exercise on the CNS in subjects with hemiparesis have also been studied to better delineate its potential impact on neural mechanisms underlying hemiparetic gait. One approach has been examining the short-term effects of submaximal TM walking on the corticospinal responses of leg muscles. In subjects with chronic stroke, changes in quadriceps excitability have been elicited with short-term exposure to self-selected TM walking [50]. Two groups of subjects with chronic hemiparesis, one that trained for 6 months in a T-AEX program and the other that was untrained, were tested with TMS before and immediately after 20 minutes of self-selected, comfortable pace TM walking. The trained group exhibited increased MEP amplitudes in paretic quadriceps, whereas the untrained group showed no change (Figure 4). In a separate study of untrained subjects with hemiparesis, this protocol was extended to include a second session of dose-time-matched stretching exercises for comparison of excitability responses with stretching versus TM walking [67]. The results of the cross-sectional study were replicated, because the submaximal TM walking had no significant effect on paretic MEP latencies or amplitudes, although the amplitudes tended to decrease in both legs. However, stretching elicited significantly larger nonparetic MEP amplitudes but with no change on the paretic side. This finding suggested that sensorimotor stimulation from stretching may have increased excitability in the former, with the possibility that longer or more intensive stretching could lead to a similar effect in the latter.

From these studies in nondisabled subjects and those with stroke, considerable evidence now exists that cortical and cortico-spinal control of the lower limbs and gait is modifiable in a short-term, transient manner. Whether such neurophysiological changes presage CNS plasticity as a viable target for long-term therapies remains to be seen. In the next section we review early results from studies that combine noninvasive measures of CNS activity associated with altered gait function.

**DURABLE LOWER-LIMB PLASTICITY AFTER STROKE**

Brain plasticity occurs with motor recovery after stroke. Longitudinal imaging and TMS mapping studies clearly show that de novo sites of brain activation become established in perilesional regions, as well as more remote areas of cortex and subcortical structures.

Upper-limb studies suggest that the lesioned hemisphere can affect cortico-muscular pathways, as repetitive TMS of the dominant, affected (but not the nondominant, unaffected) hemisphere impairs motor function to the affected hand [68]. One mechanism that may explain this control of the perilesional cortex is continued use of the affected limb, which may help maintain viable networks in the injured cortex [69–70]. Ipsilesional cortical activation has been shown to be a feature of locomotor recovery without specific training regimens [71]. Also, it is possible for cortical injury to prompt formation of axon projections

![Figure 4](image-url)

Examples of 10 averaged transcranial magnetic stimulation-induced MEPs at vastus medialis before and after single session of treadmill (TM) walking exercise: (a) trained subject’s nonparetic (NP) and paretic (P) responses and (b) responses of untrained subject. Arrows denote stimulus onset. P = paretic side, S12 = subject 12, S50 = subject 50. **Source:** Reprinted by permission from Forrester LW, Hanley DF, Macko RF. Effects of treadmill training on transcranial magnetic stimulation-induced excitability to quadriceps after stroke. Arch Phys Med Rehabil. 2006; 87(2):229–34. [PMID: 16442977]
to other cortical areas, which may promote reorganization via remodeled connections to cortical and subcortical structures [72–74].

The resultant patterns of brain reorganization after stroke appear to be strongly influenced by lesion location. Using functional magnetic resonance imaging (fMRI) techniques to study brain activations during knee movements, Luft et al. found differences in regional activations of the paretic limb versus the nonparetic limb in subjects with stroke and compared with nondisabled controls [75]. As seen with the upper limb [76], these analyses demonstrate heterogeneous CNS reorganization for lower-limb control that correlates to lesion location (Figures 5–6). Specifically, paretic knee motor control differed among survivors of stroke, such that subcortical strokes did not shift the locus of control away from M1, whereas cortical lesions induced shifts to more perilesional and contralateral control sites. Relationships to better walking function also varied by lesion location. Faster walking among subjects with brain stem lesions required lower ipsilesional M1 activity, whereas in subjects with subcortical strokes faster walking was linked to more activity in the contralesional versus ipsilesional SMC. For those with cortical lesions, faster walking was associated with increased activation in more widely distributed areas bilaterally, possibly signifying that greater compensations after cortical injury lead to better functional outcome. Future studies are needed with larger sample sizes to better define the possible links between severity of functional deficits and lesion location and whether they will indicate different rehabilitation strategies to optimize plasticity and locomotor function.

A key question then, given the apparent adaptability of the brain for lower-limb control after stroke, is whether and/or how this process can be exploited to the individual’s advantage for regaining independent mobility. Added context for the recovery of gait function is provided by a study of lower-limb EMG timing patterns to assess possible changes in motor control of hemiparetic walking after 10 weeks of physical and occupational therapies in the subacute phase poststroke [77]. While significant improvements were reported in measures of gait function, including walking velocity and indices of walking independence, no changes in EMG patterns were observed in TM tests performed at the same velocity at all time points. This finding suggests that the neuromotor control of the lower limb during walking was not reorganized by the usual therapies, which concentrated on use of the paretic limbs in the performance of gross motor skills and neurodevelopmental approaches. While Den Otter et al. concluded that locomotor functional gains could be elicited without concomitant changes in lower-limb muscle activity patterns, the results also suggest that task-specificity of practice may be a precondition to altering the underlying motor control. The results also raise questions about whether the concentration of practice in the
Figure 6.
For (a) brain stem, (b) cortical, (c) subcortical, and (d) nondisabled control subjects, activation patterns of paretic (red-yellow), nonparetic (blue), and nondisabled control knee movement (green) are superimposed onto averaged anatomical templates. Image data of subjects with left-sided stroke are flipped about midsagittal plane so that lesioned hemisphere is always on right. (d) For nondisabled control subjects, activation patterns of left- and right-sided knee movement were averaged (after appropriate flipping so that moving limb is on left). Whereas during paretic limb movement, (c) subjects with subcortical stroke and, to lesser degree, (a) brain stem subjects recruited sensorimotor cortex and supplementary motor area bilaterally, (b) almost no cortical activation is observed in subjects with cortical stroke. For nonparetic limb movement, consistent contralateral primary motor cortex activation is seen in all groups, but also markedly different from control. L = left side, R = right side. Source: Reprinted by permission from Luft AR, Forrester L, Macko RF, McCombe-Waller S, Whitall J, Villagra F, Hanley DF. Brain activation of lower extremity movement in chronically impaired stroke survivors. Neuroimage. 2005;26(1):184–94. [PMID: 15862218]
usual therapy sessions was sufficient to promote adaptations due to motor learning [77].

To date, few neuroimaging studies exist of brain activation responses secondary to sustained intensive training of lower-limb motor function. However, evidence suggests that sufficient motor practice can alter CNS control of the lower limb and gait. For example, a case study by Carey et al. used fMRI to show the feasibility of promoting brain plasticity and durable functional benefits from visuomotor training of the paretic ankle [78]. The subject was trained to use a visual tracking system to monitor volitional dorsiflexions of the paretic ankle during fMRI scans. After 16 sessions over a 4-week period, brain activation increased significantly, along with observed improvements in walking and ankle movements. Although these motor improvements were within the criterion difference of 2 standard deviations away from the baseline means, they were retained 4 months following completion of training.

Another fMRI study with four chronic survivors of stroke examined responses in cortical activity associated with ankle dorsiflexion control and lower-limb function during and after a 10-week program of PBWS TM training [79]. Serial fMRI tests were conducted at 2-week intervals, as were lower-limb Fugl-Meyer scores and walking velocity through 8 weeks of the protocol. The training produced increased activation areas in S1 and M1 regions, while functional performances improved. As function plateaued, the fMRI signals declined, a possible early indicator of learning consolidation.

Added perspective on the effects of PBWS on the CNS is gained from consideration of locomotor therapy in patients with incomplete SCI. Winchester et al. found that subjects with motor-incomplete SCI improved overground walking function that was associated with increased SMC and cerebellar activity after 12 weeks of PBWS TM training on the Lokomat robotic orthosis system [80]. Using intermuscular EMG coherence measures and TMS, Norton and Gorassini showed that training responses of incomplete SCI patients after 4 months of PBWS TM training depended on the extent of spared efferent pathways to the lower limbs [81]. The responders showed improved corticospinal connectivity in terms of increased EMG coherence at frequencies mediated by supraspinal inputs, as well as increased TMS MEP responses in the same muscles. For stroke, these improvements suggest that the degree of injury to descending pathways may have a significant effect on the capacity for the CNS plasticity to alter locomotor function, even with long-term training.

Miyai et al. conducted an intervention to study the effects of inpatient rehabilitation on eight patients who had not regained ambulatory function after 2 to 3 months of usual therapies following stroke [82]. Cortical activity was measured with NIRS during a standardized TM walking test conducted before and after a 2-month intervention that was based on a multidisciplinary neurodevelopmental approach. The regional activity changes detected from pre- and post-NIRS scans of subjects while walking showed improved symmetry in the medial primary SMCs from increased activation in the lesioned hemisphere and a reduction in the nonlesioned hemisphere. This finding parallels patterns of shifting cortical activation from nonlesioned to lesioned hemispheres in some studies of upper-limb recovery [83–84]. The change in the SMC laterality index also was significantly correlated to improved swing-phase symmetry during the posttherapy walking trials. Other activation gains were seen in the lesioned side premotor area, whereas changes in laterality of the premotor and SMAs were not significant. Perhaps the most intriguing aspect of this study was that the adaptations in CNS locomotor control resulted from interventions that were not explicitly related to gait. While the plasticity of central neural control is evident, we cannot discern the relative contributions of ongoing recovery and the therapeutic intervention.

More recent evidence suggests that an intensive practice and training regimen of T-AEX training does modify brain areas controlling the paretic leg. A preliminary report on the effects of 6 months of T-AEX training on brain maps indicates strongly that subcortical structures, including bilateral red nucleus, represent new sites of paretic knee activation using the same fMRI knee protocols [85]. Correlations between changes in both voxel-based and region of interest analyses to changes in gait peak effort walking velocity appear to support functional relevance to the new areas of activity. If pending randomized controlled trial results confirm this effect, it will suggest that extensive massed practice on the TM may stimulate motor learning and foster new or reactivate unused bilateral pathways to mediate changes in gait, with the brain stem regions assuming a prominent role in remodeling the neuromotor coupling process.

**CONCLUSIONS**

The recent advances in motor-learning-based therapies have opened new possibilities for recovery of motor
functions after stroke. The biology of central neural plasticity has emerged as a prime mechanism that may be exploited to optimize therapy for hemiparesis in the lower limb. In the area of gait rehabilitation, various methods of TM training have effectively improved walking function among individuals with hemiparesis following stroke. That such improvements can be achieved long after the expected time window for natural recovery supports the idea that the TM stimulus can promote motor learning and neuroplasticity of the lower limb. Considerable evidence now exists that supraspinal activity in the CNS, including interconnections among cortical, subcortical, and cerebellar pathways, plays a significant role in the control of lower-limb movements and gait. These direct neurophysiological findings complement imaging studies showing that natural recovery with standard therapy does foster CNS plasticity of lower-limb motor control, similar to that reported for the upper limb [75].

Although stroke often affects motor function via injured supraspinal circuitry, lesion location and size have varied effects on lower-limb function including gait, especially if the pathways from the SMC leading to and through the reticulospinal areas are selectively affected. Deficient supraspinal input to the descending tracts may cause maladaptive plasticity within the spinal level circuitry. Stroke-induced loss of cortical inhibition over spinal reflex circuits can lead to a range of “upper motor neuron” signs, including clonus, positive extensor plantar reflexes, and spasticity. Sheean suggests that these signs are due to gradual and detrimental plasticity within the spinal cord, because these changes do not appear immediately after stroke [86]. Although individual responses to TM or other motor-learning-based rehabilitation programs (e.g., robotics) may differ according to level and size of infarct, no studies to date have reported distinctions in training efficacy related to these anatomical variables and whether or to what degree spinal centers would adapt.

We hypothesize that TM or other similar locomotor training that evokes functional improvement in gait through massed practice and goal-based progressions is likely to encourage positive rather than negative adaptation at the subcortical and/or spinal levels. Our recent results show that increased peak walking velocities are associated with new subcortical and cerebellar areas becoming active in paretic knee control [85]. Although individual responses to TM or other motor-learning-based rehabilitation programs (e.g., robotics) may differ according to level and size of infarct, this finding suggests that locomotor training may not need to be overly specific to foster benefits across different stroke subtypes. That said, we are far from having the means to determine whether a given survivor of stroke is a good or bad candidate for TM therapy, as substantial differences exist in gait deficit severity among survivors of stroke [40]. More rigorous study of factors such as lesion location, size, and the associated deficit profiles are needed to develop a sound clinical basis for prescribing and implementing individualized rehabilitation programs.

In line with established models of CNS plasticity, solid indications exist that the neuroplasticity associated with improved locomotor control may be facilitated if (1) the paretic leg is actively engaged in movement practice, (2) the practice includes high volumes of repetition, and (3) the practiced movements are task-relevant with an element of problem solving (e.g., focusing on specific elements of paretic leg stepping). TM training that progresses the performance demands or goals would seem to meet these criteria, whether through gradual reductions in PBWS or increasing practice workloads through longer duration and faster velocity or immediately transferring TM practice patterns to overground walking. Evidence of short-term adaptations in lower-limb neurophysiology support the possibility of modifying the neural control of hemiparetic gait through such training regimens. However, our understanding of how to take advantage of this process is extremely limited, with most attention given to varying the structure of locomotor practice.

Critical to these suggestions is determining the generalizability of TM training after stroke. We do not know how soon after stroke TM therapy should be started to optimize responses for lasting and clinically meaningful improvements in gait function. While we do know that individuals with disparate lesion locations and severity of locomotor deficits can benefit from TM exercise training, we still have very limited knowledge on how to tailor programs to specific cases. Current studies are looking at the relative effects of duration-based versus velocity-based approaches to TM training progressions on functional outcomes and CNS plasticity. On a broader note, and regardless of what technological advances eventually become efficacious, we now know that an opportunity exists to change the course of lower-limb recovery after hemiparetic stroke. Further studies are needed to determine optimal motor-learning strategies and dose intensities to improve mobility function poststroke.
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