Neural coupling between upper and lower limbs during recumbent stepping

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Huang, Helen J., and Daniel P. Ferris. Neural coupling between upper and lower limbs during recumbent stepping. J Appl Physiol 97: 1299–1308, 2004. First published June 4, 2004; 10.1152/japplphysiol.01350.2003.—During gait rehabilitation, therapists or robotic devices often supply physical assistance to a patient’s lower limbs to aid stepping. The expensive equipment and intensive manual labor required for these therapies limit their availability to patients. One alternative solution is to design devices where patients could use their upper limbs to provide physical assistance to their lower limbs (i.e., self-assistance). To explore potential neural effects of coupling upper and lower limbs, we investigated neuromuscular recruitment during self-driven and externally driven lower limb motion. Healthy subjects exercised on a recumbent stepper using different combinations of upper and lower limb exertions. The recumbent stepper mechanically coupled the upper and lower limbs, allowing users to drive the stepping motion with upper and/or lower limbs. We instructed subjects to step with 1) active upper and lower limbs at an easy resistance level (active arms and legs); 2) active upper limbs and relaxed lower limbs at easy, medium, and hard resistance levels (self-driven); and 3) relaxed upper and lower limbs while another person drove the stepping motion (externally driven). We recorded surface electromyography (EMG) from six lower limb muscles. Self-driven EMG amplitudes were always higher than externally driven EMG amplitudes (P < 0.05). As resistance and upper limb exertion increased, self-driven EMG amplitudes also increased. EMG bursts during self-driven and active arms and legs stepping occurred at similar times. These results indicate that active upper limb movement increases neuromuscular activation of the lower limbs during cyclic stepping motions. Neurologically impaired humans that actively engage their upper limbs during gait rehabilitation may increase neuromuscular activation and enhance activity-dependent plasticity.

The purpose of this study was to determine whether evidence of neural coupling between the upper limbs and lower limbs exists in healthy subjects during a simplified stepping task. To constrain movement kinematics, we studied subjects as they exercised on a recumbent stepper that mechanically coupled upper and lower limb motion. We compared muscle activation during self-driven and externally driven stepping motions. Self-driven refers to an individual using only his/her upper limbs to drive his/her relaxed lower limbs through the stepping motion. Externally driven refers to a separate individual driving the subject’s relaxed upper and lower limbs through the stepping motion. We hypothesized that self-driven stepping motion will result in greater lower limb muscle activity compared with exter-

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nally driven stepping motion. We based this hypothesis on evidence that the upper and lower limbs of humans appear to be neurally coupled during locomotor movements (5, 14, 15, 20, 23, 48, 55, 56).

METHODS

Subjects

Twenty-four healthy subjects (14 men and 10 women; age range 19–29 yr) participated in this study. The University of Michigan Medical School Institutional Review Board granted approval for this project, and all subjects gave informed, written consent.

Recumbent Stepper

The NuStep recumbent stepper (TRS 4000, NuStep, Ann Arbor, MI) is a commercially available exercise machine that is a hybrid between a recumbent bicycle and a stair stepper. It has two handles and two pedals that are all coupled to one another, allowing users to exercise with upper and/or lower limbs (Fig. 1). The right handle and left foot pedal move together, as does the left handle and right foot pedal. The contralateral handle-pedal pairs are 180° out of phase with each other so that when the right upper limb and left lower limb are fully extended, the left upper limb and right lower limb are fully flexed. This mechanical coupling results in a simplified one degree of freedom stepping motion that preserves normal walking interlimb phase relationships. The recumbent stepper also has 10 unitless levels of resistance, created via an eddy-current disk brake. A mechanical lever allows the user to move an array of magnets closer and farther from the spinning conductive disk, thus increasing or decreasing, respectively, the resistive forces.

General Procedure

Subjects adjusted the stepper’s handle and seat positions so that their knees did not reach full knee extension. Velcro straps secured the subject’s feet to the stepper’s pedals and the subject’s torso to the stepper’s seat. A digital metronome set to a frequency of 1.25 Hz (75 beats/min) helped subjects maintain a constant stepping frequency. We recorded electromyography (EMG) from six muscles on each lower limb (Konigsberg Instruments, Pasadena, CA). We measured the pedal force for each foot during stepping using single-axis compression load cells (LCWD-1000, Omegadyne, Sunbury, OH). We also measured joint angles using electrogoniometers (Biometrics, Ladysmith, VA). All subjects were given time to practice stepping using each combination of upper and lower limb exertions. For each trial, we collected data for 10 s after subjects verbally confirmed that they were comfortable with the test condition. We recorded two trials for each test condition and randomized trial order for each subject to minimize potential learning effects. Subjects also rested for ~20 s between trials and had the option to rest longer.

Experimental Design

We conducted two separate experiments. In the first experiment, the upper and lower limb motions were mechanically coupled as dictated by the stepper’s design. In the second experiment, we used two recumbent steppers to test a similar protocol but with the upper and lower limb kinetics decoupled. In this second experiment, we placed one recumbent stepper behind another and modified the front stepper with long handle extensions (Fig. 2). This allowed subjects to drive the front stepper with their upper limbs while sitting in the back stepper. Decoupling the upper and lower limb kinetics in this manner prevented subjects from pushing with their lower limbs to lessen the work performed by the upper limbs.

Experiment 1. Subjects (n = 20) stepped using four different combinations of upper and lower limb exertions. We classified these exertions as “active” and “passive” in reference to the subject’s conscious intent. There was one condition in which subjects stepped using both their upper and lower limbs (active arms and legs). This condition provided us with information about the normal timing of lower limb muscle activation during active recumbent stepping. There were three conditions in which subjects attempted to totally relax their lower limbs (i.e., passive lower limbs). For the simplest passive condition, subjects rested their hands on their laps as another person drove their lower limbs through the stepping motion (externally driven). For a second passive condition, subjects’ hands and feet were fixed to the handles and pedals using Velcro mitts and straps, respectively, as another person drove their relaxed upper and lower limbs through the stepping motion (externally driven arms and legs). We tested this condition to determine whether passive movement of the upper limbs made a difference in the results. For the last passive condition, subjects actively used their arms to drive their relaxed lower limbs through the stepping motion (self-driven).
We examined three levels of resistance for the self-driven passive condition: easy, medium, and hard. These resistance levels were determined before testing for each subject. Each subject's hard resistance level corresponded to the maximum resistance that he/she could maintain using just his/her arms moving at the designated frequency for at least 20 s. The medium resistance level corresponded to ~60% of the subject's hard resistance level. The easy resistance level was the lowest resistance setting on the recumbent stepper. Because the level of resistance was not important for the externally driven conditions, the resistance was set at easy for all externally driven trials.

Experiment 2. The main difference between this experiment and the first experiment was that a second recumbent stepper was used to decouple the kinetics of the pedals and handles (Fig. 2). We tested four healthy male subjects under three conditions: 1) externally driven, 2) upper limb exertion (easy), and 3) upper limb exertion (hard). In the externally driven condition, subjects were instructed to attempt to totally relax their upper and lower limbs, as other people drove the subject's upper and lower limbs through the range of motion at the designated frequency. In the upper limb exertion conditions, subjects were instructed to actively use their upper limbs to drive the front stepper at the designated frequency while relaxing their lower limbs as another person drove their legs through the motion. We instructed the person driving the subject's lower limbs to match the subject's upper limb movement. For the upper limb exertion conditions, we examined two resistance levels: easy and hard. These resistance levels were determined before testing for each subject. Each subject's hard resistance level corresponded to the maximum resistance that he/she could step at using just his/her arms while maintaining the designated frequency for at least 20 s.

Data Acquisition and Analysis

In the first experiment, we collected EMG, joint angle (hip, knee, and ankle), and pedal force data during each trial. In the second experiment, we collected EMG and joint angle (knee and elbow) data during each trial. For both experiments, a personal computer sampled each data channel at 1,000 Hz via an analog-to-digital board.

EMG. For experiments 1 and 2, we measured muscle activity from six muscles on each lower limb [vastus lateralis (VL), vastus medialis (VM), medial hamstrings (MH), medial gastrocnemius (MG), tibialis anterior (TA), and soleus (SO) muscles] using surface electrodes (Konigsberg Instruments). For each electrode site, we shaved and cleaned the area with alcohol. We then placed EMG surface electrodes (diameter, 1.1 cm; interelectrode distance, 3.5 cm) in the middle of the muscle belly along its long axis. For the SO, however, we placed the electrodes over the lateral distal third of the muscle belly rather than the center to avoid potential cross talk from the gastrocnemius. We then visually examined each EMG signal for noise and cross talk (52).

We placed tape over all electrodes and covered the subject’s lower limbs with elastic stockings to minimize mechanical artifacts. To compare muscle activation across conditions, we calculated the averaged normalized root mean square (RMS) EMG amplitude for each condition and muscle for flexion and extension phases. We processed EMG with a fourth-order high-pass Butterworth filter with zero phase shift (cutoff frequency of 20 Hz) and then full-wave rectified the signals. For each phase, condition, muscle, and subject, we averaged RMS amplitudes for six step cycles. We then normalized the averaged RMS amplitudes to the maximum RMS amplitude calculated for each muscle across conditions for each subject to reduce intersubject variability (54).

To compare EMG timing across conditions, we used cross-correlation analysis to calculate R values and lag times.

Joint angles. For experiment 1, we measured joint angles of the hip, knee, and ankle on both lower limbs using twin-axis electromagnetic goniometers (Biometrics), placed along the sagittal plane. For experiment 2, we measured knee joint angles on both lower limbs and elbow joint angles on both upper limbs using twin-axis electrogoniometers (Biometrics) placed along the sagittal plane. Electrogoniometers were zeroed as subjects stood in a neutral position. From these data, we identified the beginning and end of step cycles. We also used goniometer data to define flexion and extension phases.

Pedal forces. For experiment 1, we measured the pedal force for each foot during stepping using a set of three compression load cells (LCWD-1000, Omegadyne). These load cells were coplanar, positioned in a tripod formation between two custom-shaped aluminum plates for each foot. The summation of the three load cells for each foot provided a measure of the force between the foot and pedal. Before each data collection, we calibrated each load cell within the aluminum plates with known weights. We summed the three load cell measurements for each foot plate and then filtered the data with a fourth-order low-pass Butterworth filter with zero phase shift (cutoff frequency of 6 Hz). We calculated the mean forces during lower limb flexion and extension for six step cycles per condition. We then normalized the forces to the maximum mean force across conditions for each subject. Last, we averaged the forces across all subjects for the flexion and extension phases. We were only able to collect pedal forces on 10 subjects.

Statistical Analyses

For each experiment, we used a repeated-measures ANOVA to determine whether there were significant differences across conditions. For the ANOVA analyses of EMG in the six muscles, we set P < 0.0083 for the significance level (Bonferroni correction). When the ANOVA indicated a significant difference, we used Tukey’s honestly significant difference (THSD) post hoc test to determine which conditions were significantly different (P < 0.05).

RESULTS

Experiment 1: One NuStep with Upper and Lower Limb Kinetics Coupled

Self-driven stepping motion resulted in greater lower limb muscle activation compared with externally driven stepping motion. Externally driven EMG raw data showed little EMG activity for all six muscles, whereas distinctive rhythmic EMG bursts were visible in self-driven (hard) EMG raw data (Fig. 3). Group-averaged EMG profiles indicated that lower limb muscle activation for the self-driven (hard) condition had similar patterns compared with the active arms and legs condition (Fig. 4). Onset of the EMG bursts during self-driven (hard) and active arms and legs conditions occurred at nearly the same time in the step cycle. Lag times at maximum correlation between self-driven (hard) and active arms and legs were close to zero (Table 1). This confirmed that self-driven (hard) and active arms and legs EMG bursts had similar timing. Cross-correlation results comparing medium and easy self-driven conditions to active arms and legs also showed similar timing in muscle activation (not shown).

For all six muscles during both flexion and extension, the normalized EMG amplitudes for the three self-driven conditions were greater than the amplitudes for the externally driven and externally driven arms and legs conditions (Fig. 5). Across all six muscles, self-driven EMG amplitudes for the flexion phase were ~35, 42, and 55% for easy, medium, and hard resistances, respectively. In contrast, externally driven and externally driven arms and legs conditions had EMG amplitudes of 30% across all six muscles. Similarly, self-driven EMG amplitudes averaged for all six muscles for the extension phase were 38, 46, and 55% for easy, medium, and hard resistances, respectively. Externally driven and externally driven arms and legs conditions had EMG amplitudes of 32 and 31%, respectively, averaged for all six muscles. Statistical
analyses indicated that, for both flexion and extension phases, self-driven (hard) and active arms and legs were significantly different (THSD, \( P < 0.05 \)) from externally driven and externally driven arms and legs conditions for a majority of the muscles (Fig. 5).

During self-driven stepping, greater upper limb exertion led to greater lower limb muscle activation. Self-driven (easy), self-driven (medium), and self-driven (hard) EMG amplitudes increased in a stepwise manner (Fig. 5). Self-driven (medium) was not significantly different from self-driven (easy) for any of the muscles (THSD, \( P > 0.05 \)). Self-driven (hard) was significantly different from self-driven (easy) for the MH, SO, and TA muscles during flexion and for the VL, VM, MH, SO, and TA during extension (THSD, \( P < 0.05 \)). Self-driven (hard) was significantly different from self-driven (medium) for the TA and SO muscles during flexion (THSD, \( P < 0.05 \)).

Pedal force profiles were similar for self-driven and active arms and legs stepping (Fig. 6A). Both conditions had higher peak forces during the extension phase than during the flexion phase, and the two conditions demonstrated similar timing of

<table>
<thead>
<tr>
<th>Muscle</th>
<th>( R ) Value</th>
<th>Lag Time, % cycle</th>
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<tbody>
<tr>
<td>VL</td>
<td>0.84±0.10</td>
<td>-1±11</td>
</tr>
<tr>
<td>VM</td>
<td>0.84±0.09</td>
<td>-3±14</td>
</tr>
<tr>
<td>MH</td>
<td>0.89±0.06</td>
<td>4±12</td>
</tr>
<tr>
<td>MG</td>
<td>0.90±0.06</td>
<td>1±5</td>
</tr>
<tr>
<td>SO</td>
<td>0.90±0.06</td>
<td>-2±13</td>
</tr>
<tr>
<td>TA</td>
<td>0.88±0.10</td>
<td>2±14</td>
</tr>
</tbody>
</table>

Values are means ± SE. A positive lag time indicates that self-driven (hard) follows active arms and legs. A negative lag time means that self-driven (hard) begins before active arms and legs. Mean lag times for all muscles are <4% of a step cycle from zero, indicating that self-driven (hard) and active arms and legs have similar timing. EMG, electromyography; VL, vastus lateralis; VM, vastus medialis; MH, medial hamstrings; MG, medial gastrocnemius; SO, soleus; TA, tibialis anterior.
force increase during the start of the extension phase. In contrast, the externally driven pedal force profile had a different pattern (Fig. 6A). Overall, the externally driven condition had lower peak forces during the extension phase than during the flexion phase. There was a slight rise in pedal force at the end of extension and the beginning of flexion to decelerate the lower limb. Self-driven mean flexion pedal forces were 45, 42, and 40% for easy, medium, and hard resistances, respectively (Fig. 6B). Externally driven mean flexion pedal forces were 50% for both conditions (externally driven, externally driven arms and legs). For the extension phase, self-driven mean pedal forces were 39, 44, and 53% for easy, medium, and hard conditions, respectively (Fig. 6C). Both externally driven and externally driven arms and legs mean extension pedal forces were 41%. Self-driven and externally driven mean pedal forces were not significantly different from each other for either phase (THSD, \( P > 0.05 \)). Active arms and legs mean extension force (93%) was significantly different from externally driven mean extension force (THSD, \( P < 0.05 \)).

Joint angle data indicated that ranges of motion for the hip, knee, and ankle were similar for each subject across all conditions. Hip, knee, and ankle profiles during recumbent stepping were regular and synchronized, having a triangular pattern (Fig. 3). The average hip angle excursion for all subjects was 73–43° of hip flexion (for all 3 joints, 0° was standing posture). The average knee angle excursion was 75–23° of knee flexion. The average ankle angle excursion was 19° of dorsiflexion to −3° of plantarflexion. There was no significant difference in range of motion between conditions (ANOVA, \( P > 0.05 \)).

Fig. 5. Averaged normalized root mean square (RMS) EMG with standard error bars for all subjects for flexion and extension phases in experiment 1. White columns represent externally driven conditions, whereas gray columns indicate self-driven conditions. Black columns correspond to the active condition. *Significantly different from the externally driven condition on the far left [Tukey’s honestly significant difference (THSD), \( P < 0.05 \)]. For both flexion and extension, active arms and legs and self-driven (hard) RMS EMG amplitudes were significantly different from both externally driven conditions for a majority of the muscles (THSD, \( P < 0.05 \)). Externally driven and externally driven arms and legs RMS EMG amplitudes were not significantly different for any of the 6 muscles in either phase (THSD, \( P > 0.05 \)).
condition (Figs. 7 and 8). Group-averaged EMG showed that the coupled and decoupled protocols resulted in similar EMG patterns (Fig. 7). Externally driven EMG profiles were fairly constant with no distinct bursts in any of the muscles. Upper limb exertion (hard) EMG profiles had distinct EMG bursts for the VL, VM, and TA muscles. Both upper limb exertion conditions had greater normalized EMG RMS amplitudes compared with the externally driven condition for both flexion and extension phases (Fig. 8). Statistical analyses indicated that upper limb exertion (hard) was significantly different (THSD, \(P < 0.05\)) from externally driven for a majority of the muscles during both flexion and extension phases.

**DISCUSSION**

Our main finding is that rhythmic upper limb activity increases lower limb neuromuscular recruitment when healthy subjects attempt to relax their lower limbs. In experiment 1, self-driven conditions had greater EMG amplitudes compared with externally driven and externally driven arms and legs conditions. In experiment 2, upper limb exertion conditions had greater EMG amplitudes compared with the externally driven condition. Thus upper limb activity resulted in greater lower limb muscle activity regardless of whether upper limb and lower limb kinetics were coupled or decoupled. An important aspect of this finding was that lower limb muscle activation during the self-driven conditions was dependent on upper limb effort. As resistance level increased for the upper limbs, EMG amplitudes for the lower limbs muscles also increased.

At all resistance levels, the timing of the self-driven EMG bursts in the lower limbs was similar to the timing of active stepping EMG bursts. Muscles were primarily active during the concentric phase of the stepping motion. For the VL, VM, and SO, this was the extension phase. For the TA, this was the flexion phase. The biarticular MH and MG showed some activation during both flexion and extension phases. The concentric timing of the uniarticular muscles indicates that the increased muscle activation during self-driven stepping was probably not a result of stretch reflexes. If the stretch reflexes had caused the muscle activation, the EMG bursts would have occurred during and/or slightly after the muscles were stretched in their eccentric phase.

One potential neural mechanism responsible for lower limb muscle activation during self-driven stepping is spinal connections in locomotor neural networks. Humans have spinal connections coordinating left and right lower limbs during stepping (19) in a manner similar to cats (18). Cats also have spinal connections between cervical and lumbar locomotor networks coordinating forelimb and hindlimb muscle activation patterns (27, 31, 41). Indirect evidence indicates that humans also coordinate upper limb and lower limb movements during locomotion through spinal connections, reviewed in Refs. 14 and 56. Humans demonstrate natural frequency locking between upper limb movement and lower limb movement during walking, crawling, and swimming (48). Mechanical and electrical perturbations to the lower limbs result in short-latency reflex responses in the upper limbs during walking but not during standing or sitting (15, 23). Furthermore, near-infrared spectroscopic topography reveals that there is little brain activation in presumed arm areas of primary sensorimotor cortex during human walking (32). In contrast, active arm swinging

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**Experiment 2: Two NuSteps with Upper and Lower Limb Kinetics Decoupled**

Upper limb exertion conditions resulted in greater lower limb muscle activation compared with the externally driven
during standing activated cortical areas not activated during gait (lateral and rostral parts of primary sensorimotor cortex) (32). These findings point to humans having similar spinal interlimb connections as cats and other quadruped vertebrates.

A second potential neural mechanism that could explain our findings is neural cross talk above the spinal cord level. When humans perform muscle contractions greater than a certain force threshold, unintended muscle activity frequently occurs in other contralateral and ipsilateral muscles (1, 2, 10, 17, 21, 57). When the unintended muscle activity occurs in contralateral homologous muscles, it has traditionally been referred to as mirror movements or contralateral irradiation. Several observations suggest a cortical origin of the unintended activation, potentially related to a decrease in inhibition along the corpus callosum (29, 42). Other studies have found evidence for a different type of neural cross talk that occurs when humans perform rhythmic upper limb movements combined with rhythmic ipsilateral lower limb movements (3, 9, 28, 44), reviewed in Ref. 43. These studies indicate that the nervous system has a natural tendency to prefer in-phase movements of ipsilateral limbs rather than out-of-phase movements.

These observations of supraspinal neural cross talk do not appear to be relevant to our findings, however. The lower limb muscle activation during self-driven stepping had a timing pattern similar to active stepping (ipsilaterally out of phase). Thus the unintended lower limb muscle activity did not occur at the same time as ipsilateral upper limb muscle activity. Both of the neural cross-talk forms identified above would be associated with concurrent activation of upper and lower limb muscles. On the basis of the timing of the lower limb muscle activity, it seems likely that the neural signal responsible for activating lower limb motoneurons was filtered through neural networks responsible for active stepping.

The TA demonstrated the greatest EMG amplitudes of all the muscles during self-driven conditions. The mean TA EMG during self-driven (hard) was even higher than during the active-stepping condition. It is possible that the subjects’ strong intent to inhibit lower limb muscle activation may have decreased inhibition onto TA motoneurons via reciprocal inhibition from the plantarflexors (12, 13, 35, 36). This is only speculation, however, and future electrophysiological studies should yield more insight.

A limitation of this study was the necessity to rely on subjects’ intent to not use their lower limbs during the self-driven conditions. Subjects did not have prior knowledge of the study purpose or hypothesis. We instructed all subjects to completely relax their lower limbs before each trial collection. Subjects verbally confirmed that they understood the instructions and did not intentionally use their lower limbs to push the pedals. They may have been tempted to cheat if fatigued, but our protocol limited that possibility because of the short data-collection periods and frequent rest breaks. On average, there was also ~20 s of rest between each 15-s trial. All trials were randomized, reducing the probability that subjects would have to work actively for extended periods of time. Furthermore, we analyzed step cycles from the middle of the data-collection period, when subjects were most likely not fatigued. Most importantly, the results from experiment 2 indicate that increased lower limb recruitment still occurs with upper limb exertion even when pedals and handles have been kinetically decoupled. Cheating in experiment 2 could not decrease the work performed by the upper limbs. We conclude that any muscle activity during self-driven conditions was indeed unintentional.

Regardless of mechanism, the apparent neural coupling between upper limbs and lower limbs during cyclic stepping movements could be useful during rehabilitation. We only tested subjects using one exercise device (i.e., a recumbent stepper), but it seems feasible that the neural coupling effect would also occur during other types of rhythmic upper and
lower limb movements (e.g., upper and lower limb cycling). If neurologically impaired subjects demonstrate similar neural coupling, repetitive upper and lower limb motor tasks may help improve muscle coordination compared with lower limb motor tasks alone. The specificity of learning hypothesis (40) implies that the upper and lower limb motor task should be as close to possible normal walking to provide the greatest functional benefit for overground locomotion. However, locomotor neural networks appear to be active during other cyclic lower limb motor tasks such as cycling (8, 37, 45–47). A recent study on spinally transected rats (6) compared the effects of manually assisted treadmill stepping and mechanically assisted cycling. The study found that cycling produced similar or better recovery of locomotor function than treadmill stepping (6). In addition, Werner et al. (49) demonstrated that upright use of an elliptical-type trainer produces improvements in overground walking ability comparable to improvement with manually assisted treadmill stepping. These findings indicate that rhythmic stepping tasks are likely to contribute to gait rehabilitation even if they do not exactly follow the same movement dynamics as walking.

When recumbent stepping is compared with walking, there are some obvious differences between the dynamics of the two motor tasks. First, the hip joint does not hyperextend at the end of the extension phase during recumbent stepping as it does during walking. Second, the bottom of the foot is continually in contact with the pedal during lower limb flexion while recumbent stepping, unlike lower limb flexion while walking. Third, the ankle and knee joints only flex and extend once during a complete cycle in recumbent stepping but flex and extend twice during each cycle in walking. Fourth, recumbent stepping uses a reclined posture instead of the vertical posture characteristic to walking. These four differences between recumbent stepping and walking clearly produce discrepancies in sensory feedback. In particular, the lack of hip flexor muscle stretch at the end of lower limb extension and the presence of cutaneous pressure during lower limb flexion are potentially critical to gait rehabilitation (7, 16, 19, 22, 33, 34, 39, 41, 50). Future research is needed to determine how critical these discrepancies would be on neurorehabilitation.

There are some benefits to recumbent stepping that partially offset limitations in task specificity and make it appealing as an alternative and/or supplemental gait rehabilitation therapy. Self-assisted recumbent stepping offers a way to give patients control over the amount and timing of physical assistance as they practice stepping. Currently, locomotor training for gait rehabilitation relies on external assistance from therapists or robots to induce stepping in neurologically impaired patients (4, 11, 24, 25, 38, 51). Self-assisted gait rehabilitation devices eliminate these expensive manual labor costs. Additionally, self-assisted devices may provide more appropriately timed and scaled physical assistance for stepping. Another advantage of recumbent steppers or other future self-assisted gait rehabilitation devices is that they offer neurologically impaired individuals a means to practice stepping at home. Because individuals could practice whenever they choose, they could complete more training sessions compared with therapies that are only available at clinics. Performing a stepping task that is only 50% similar to walking daily may be more beneficial to gait rehabilitation than performing a stepping task that is 95% similar to walking once a week. More generally, it appears that self-assisted stepping devices of some design may be able to capitalize on neural coupling between the upper and lower limbs during rhythmic locomotor-like tasks.

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