Upper and Lower Limb Muscle Activation Is Bidirectionally and Ipsilaterally Coupled

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ABSTRACT

HUANG, H. J., and D. P. FERRIS. Upper and Lower Limb Muscle Activation Is Bidirectionally and Ipsilaterally Coupled. Med. Sci. Sports Exerc., Vol. 41, No. 9, pp. 1778–1789, 2009. Purpose: There are neural connections between the upper and lower limbs of humans that enable muscle activation in one limb pair (upper or lower) to modulate muscle activation in the other limb pair (lower or upper, respectively). The aims of this study were to extend previous findings regarding submaximal exercise to maximal effort exercise and determine whether there is an ipsilateral or contralateral bias to the neural coupling during a rhythmic locomotor-like task. Methods: We measured upper and lower limb muscle activity, joint kinematics, and limb forces in neurologically intact subjects (n = 16) as they performed recumbent stepping using different combinations of upper and lower limb efforts. Results: We found increased muscle activation in passive lower limbs during active upper limb effort compared with passive upper limb effort. Likewise, increased muscle activation in passive upper limbs occurred during active lower limb effort compared with passive lower limb effort, suggesting a bidirectional effect. Maximal muscle activation in the active lower limbs was not different between conditions with active upper limb effort and conditions with passive upper limb movement. Similarly, maximal muscle activation in the active upper limbs was not different between conditions with active lower limb effort and conditions with passive lower limb movement. Further comparisons revealed that neural coupling was primarily from active upper limb muscles to passive ipsilateral lower limb muscles. Conclusions: These findings indicate that interlimb neural coupling affects muscle recruitment during maximal effort upper and lower limb rhythmic exercise and provides insight into the architecture of the neural coupling. Key Words: INTERLIMB COUPLING, NEURAL COUPLING, EMG, COORDINATION, BILATERAL DEFICIT

Humans naturally couple limb movements. It is easier to move two limbs in the same direction rather than in the opposite directions or when using homologous muscles rather than nonhomologous muscles (2,20,27). Humans also choose coordination patterns that maintain an integral frequency ratio between the upper limbs and lower limbs during rhythmic whole-body tasks such as walking, swimming, and crawling (29). Muscle activation patterns (3,10,16) and reflex responses (4,5,9) during multijoint and/or multilimb tasks suggest that the coupled limb movements have a neural component. For example, during walking, humans use shoulder muscles to help drive backward arm swing in-phase with the backward swing of the contralateral leg. When the arms are bound to restrict arm swing, there is still muscle activation in the shoulder muscles (3). A likely candidate for facilitating coordinated interlimb movements in humans is propriospinal connections between upper limb neural networks and lower limb neural networks (7,33).

Studies have shown that adding upper limb movement or effort concurrently with lower limb movement during rhythmic tasks may improve lower limb muscle recruitment. We previously demonstrated that increased upper limb muscle activity resulted in greater muscle activation in passively moving legs of neurologically intact subjects performing recumbent stepping (15,17). Kawashima et al. (18) similarly examined the effects of resting, passive, and active arm swing on passive lower limb muscle activation during a reciprocal leg swinging task. They found that passive arm swing improved muscle activation patterns in passively moved lower limbs compared with a resting arms condition in incomplete spinal cord injured individuals.
These two studies demonstrate that upper limb activation or afferent feedback can increase or improve lower limb muscle activation.

When examining muscle recruitment during multilimb tasks, another neurally mediated interlimb effect is the so-called bilateral deficit (14). A bilateral deficit occurs when the force output of two bilateral limbs performing a task simultaneously is less than the sum of the force output of each single limb performing the same task. Deficits in muscle activation often parallel the deficits in force production (22,28) and suggest that the bilateral deficit may represent a limitation in neuromuscular activation (16). Bilateral deficits occur in both the upper and lower body during isometric tasks (19,21–24). Dynamic and/or multijoint lower limb movements also typically produce bilateral deficits (25,28,30). Despite the common use of simultaneous upper and lower limb tasks for exercise (such as using elliptical trainers), no study has examined how maximum muscle activation is affected by simultaneous upper and lower limb rhythmic exercise compared with just upper or lower limb exercise.

The general purpose of this study was to examine the effect of maximal voluntary upper limb muscle activation on lower limb muscle activation during a rhythmic task in neurologically intact humans. A secondary general purpose was to examine the same effect, but in the reverse direction (the effect of maximal voluntary lower limb muscle activation on upper limb muscle activation). We wanted to answer several specific questions. First, does maximal voluntary effort in the upper (or lower) limbs lead to increased muscle recruitment of passive lower (or upper) limbs during rhythmic movement? This question seeks to understand whether interlimb neural coupling can increase muscle recruitment of passively moving limbs. We hypothesized that active effort in the upper (or lower) limbs will increase muscle activation in passive lower (or upper) limbs. Second, does simultaneous upper and lower limb maximal effort produce more or less lower (or upper) limb muscle activation compared with only lower (or upper) limb maximal effort. This second question seeks to understand whether interlimb neural coupling can increase muscle recruitment during maximal effort. We hypothesized that simultaneous upper and lower limb maximal effort would produce greater lower limb muscle activation compared with only lower limb maximal effort. Third, does single upper (or lower) limb maximal effort produce more or less muscle activation in the ipsilateral or contralateral lower (or upper) limb? This third question seeks to understand whether interlimb neural coupling is more ipsilateral or contralateral in nature. We hypothesized that single limb effort would produce more muscle activation in the in-phase contralateral limb. To examine these questions, we studied subjects performing recumbent stepping similar to our previous works (15,17), but modified the stepping device to allow maximal effort at a constant stepping frequency (Fig. 1A).

**METHODS**

**Subjects**

Sixteen healthy female subjects (age range, 18–29 yr) participated in this study. We could only test subjects who did not exceed an upper limit of the stepper’s power output. Subjects provided informed written consent, and the University of Michigan Medical School Institutional Review Board approved the protocol and consent form. The study complied with the Declaration of Helsinki.

**Computer-Controlled Recumbent Stepper**

We have modified a commercially available recumbent stepper (TRS 4000; NuStep, Inc., Ann Arbor, MI) to have computer-controlled real-time resistance and force measuring capabilities (Fig. 1A). We used RT Lab Solo software (Opal-RT, Montréal, Québec, Canada) to customize control of the servomotor (Kollmorgen, Radford, VA) that powers the recumbent stepper. We used the position sensor in the servomotor to measure the kinematics of the recumbent stepper. The stepper has one degree of freedom and thus needs just one position sensor to describe its kinematics.

For this study, we programmed the recumbent stepper to follow a prescribed sine wave position profile (Fig. 1B) to
produce a smooth and consistent stepping at a set frequency (75 bpm, equivalent to the stepping frequency of walking at 1.25 m s⁻¹). The harder the subjects pushed or pulled, the more resistance they encountered to maintain a constant stepping frequency and minimize error. The largest errors occurred in the Active Upper & Active Lower (AU–AL) condition, reaching an average peak error of 10 ± 3% (mean ± SD) of the stepper range of motion and an average error of 5.7 ± 1.7% across the cycle. Some design limitations made it difficult to further reduce the error, but all data collected were within <1% of the target stepping frequency. To measure the force that each arm contributed to the stepping motion, we mounted a single-axis load cell (Strainsert, West Conshohocken, PA) in each handle. We also mounted a single-axis load cell (LCWD-1000; Omegadyne, Sunbury, OH) in the connecting link of each handle–pedal unit to measure the total force each contralateral upper–lower limb pair supplied to the stepping kinematics. We found that the sum of the handle and pedal torques equaled the handle–pedal unit torque. Based of this, we could determine the force contribution of each lower limb to the stepping motion.

Protocol

Subjects performed recumbent stepping using different combinations of upper (U) and lower (L) limb effort, being either active (A) or passive (P). Active effort was always with maximal effort, and passive effort was with minimal effort or as relaxed as possible. Subjects could choose any combination of pushing and pulling to drive the stepping motion, unless specifically told to just push or just pull (for the single limb conditions).

We used Velcro gloves to attach the hands to the handles and foot straps to attach the feet to the pedals during passive conditions. This adaptation allowed subjects to be as passive as possible because they did not have to actively hold the handles or keep their feet on the pedals throughout the stepping motion. A torso strap minimized torso movement during stepping. The seat position was set so that the subject’s knees were near full extension but could not lock out. The position of each handle was adjusted for subject comfort (Fig. 1A). For each condition, we collected 15 s of data, approximately six to eight complete stride cycles. The order of the conditions was pseudorandomized for each subject. There was an average of 1 min of rest between conditions.

Question 1: Does maximal effort in the upper (or lower) limbs increase muscle activation in passive lower (or upper) limbs? To answer our first question, we compared lower limb muscle activation between Passive Upper & Passive Lower (PU–PL) and Active Upper & Passive Lower (AU–PL) to determine whether maximum voluntary effort in the upper limbs leads to greater muscle activation in the passive lower limbs. We also looked at the reverse direction and compared Passive Upper & Active Lower (PU–AL) to PU–PL to determine whether maximum voluntary effort in the lower limbs leads to greater muscle activation in the passive upper limbs.

Question 2: Does simultaneous upper and lower limb maximal effort increase lower (or upper) limb maximal effort compared with only lower (or upper) limb maximal effort? To answer our second question, we compared lower limb muscle activation between AU–AL and PU–AL conditions. We also compared upper limb muscle activation between AU–AL and AU–PL conditions. These comparisons will determine whether simultaneous upper and lower limb maximal effort produces more or less muscle activation compared with only lower (or upper) limb maximal effort.

Question 3: Does single limb maximal effort increase muscle activation in the ipsilateral or contralateral limb? To answer our third question, we examined the effect of single upper limb effort on muscle activation of passive lower limbs to determine whether there was more muscle activation in the ipsilateral or contralateral passive lower limb. We tested conditions of just left upper limb effort, Active Left Upper and Passive Lower, and just right upper limb effort, Active Right Upper & Passive Lower. For these conditions, subjects were instructed to push and pull using just their left (or right) upper limb while the right (or left) upper limb was as passive as possible. We also tested conditions instructing subjects to just pull with a single upper limb, Active Left Upper Pulling & Passive Lower and Active Right Upper Pulling & Passive Lower. Similarly, we tested conditions instructing subjects to just push with a single upper limb, Active Left Upper Pushing & Passive Lower and Active Right Upper Pushing & Passive Lower. By focusing active effort of a single upper limb to just pulling (or pushing), we could potentially observe how specific upper limb muscle groups affect passive lower limb muscle activation. During these conditions, the device drove the stepping motion for the portion of the stride when the subject was instructed to be passive until reaching the portion of the stride when the subject was instructed to push (or pull) with the specified single upper limb. To examine bidirectionality, we also tested a Passive Upper & Active Left Lower and a Passive Upper & Active Right Lower condition to examine whether any ipsilateral or contralateral coupling also occurred in the lower to upper direction.

Data Acquisition and Analysis

Two computer systems sampled all of the data signals at 1000 Hz. We used a common data signal sampled in both systems to synchronize the data offline.

EMG

We measured muscle activity from 16 muscles, four muscles on each limb, using a surface EMG system with an EMG bandwidth of 20–450 Hz (Delsys, Boston, MA). On each lower limb, we measured muscle activity from the vastus medialis (VM), medial hamstrings (MH), tibialis anterior (TA), and soleus (SO). On each upper limb, we measured
muscle activity from the anterior deltoid (AD), posterior deltoid (PD), biceps brachii (BB), and triceps brachii (TB). We cleaned each electrode site with rubbing alcohol, placed the electrode sensor over the muscle belly along the long axis, and then secured the electrode with tape. To further minimize mechanical artifact, we wrapped excess loose electrode wires to the limbs with an elastic foam wrap. We processed the EMG data with a second-order high-pass Butterworth filter (cutoff frequency, 20 Hz) with zero phase lag to attenuate low-frequency components such as mechanical artifact. We then full wave-rectified the EMG data signals.

**Joint Angles**

We measured joint angles of the ankles, knees, and hips on both legs and the elbows of both arms using twin-axis

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**PASSIVE LOWER LIMB CONDITIONS**

<table>
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<tr>
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<th>AU-PL, Active Upper - Passive Lower</th>
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<tr>
<td><strong>Upper Limb EMG [μV]</strong></td>
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<tr>
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<tr>
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**Group Data**

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<td>R. Tibialis Anterior</td>
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<td>R. Soleus</td>
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**Group EMG RMS**

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<td>Lower Limb EMG RMS [%]</td>
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<td>R. Tibialis Anterior</td>
<td>100</td>
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<tr>
<td>R. Soleus</td>
<td>100</td>
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** Joint Angle [deg]**

- **Knee**
  - Flex
  - 90
  - 0
  - Ext

**Force [N]**

- **Handle**
  - 130
  - 0
  - 210
- **Pedal**
  - -180
  - 0
  - 180

**Stride**

- Pull
  - 123
  - 0
  - 210
- Push
  - -100
  - 0
  - 100

**Stride cycle [%]**

- 0
  - 50
  - 100

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**FIGURE 2**—Data from the right limbs for the passive lower limb conditions, PU–PL (black) and AU–PL (gray). Representative single-subject data (A) and group EMG mean profiles (B) show minimal EMG activity in the PU–PL condition, whereas the AU–PL condition had rhythmic bursts of EMG. Group EMG RMS amplitudes with standard error bars (C) for AU–PL were significantly greater than PU–PL for the VM, MH, TA, and SO muscles (*significantly different, AU–PL > PU–PL; THSD, P < 0.05). Dotted line equals 100%. Note different y-axes between A and B. The knee joint angle profiles and passive pedal forces were similar between the two conditions. Handle forces during the AU–PL condition had a clear increase in pulling force during the upper limb flexing (and lower limb extending) phase and pushing force during the upper limb extending (and lower limb flexing) phase.
electrogoniometers placed along the sagittal plane (Biometrics Ltd., Ladysmith, VA). The electrogoniometers were zeroed with the limbs in the anatomically neutral position. We processed the joint angle data with a second-order low-pass Butterworth filter (cutoff frequency, 6 Hz) with zero phase lag.

**Kinetics**

We calculated the forces that each hand and foot contributed to the stepping motion via single-axis load cells (Fig. 1). We measured the force exerted by each hand through a load cell mounted in the handle and the total force exerted by each contralateral hand–foot pair through a load cell mounted in a connecting link on the machine. We filtered these force data using a second-order low-pass Butterworth filter (cutoff frequency, 6 Hz) with zero phase lag. Using the measured forces and moment arm relationships, we calculated the torques associated with each handle and handle–pedal unit. To determine the pedal torque, we subtracted the handle torque from the handle–pedal unit torque. We then divided the pedal torques by the pedal moment arm to find the pedal forces.

**Calculation of Mean Profiles**

To compare EMG patterns between conditions, we calculated group EMG mean profiles over a stride cycle for each condition. The beginning and end of each stride corresponded with the left lower limb and right upper limb at full extension as indicated from the position data. We first calculated an intrasubject EMG mean profile for a stride cycle per condition. We then calculated a group EMG mean profile for each condition by averaging all of the intrasubject EMG mean profiles for that condition. We used the same procedure for the joint angle and force mean profiles.

**Calculation of EMG Root-Mean-Square**

To compare EMG amplitudes across conditions, we calculated a group-averaged (n = 16) normalized EMG root-mean-square (RMS) for each muscle and condition. The data for left and right muscles were analyzed independently. For each muscle, we calculated EMG RMS during the half of the stride when the muscle was concentrically contracting. We calculated an intrasubject average EMG RMS for each muscle per condition. We then normalized the lower limb EMG RMS amplitudes for each muscle (left VM, MH, SO, and TA; right VM, MH, SO, and TA) to the intrasubject average EMG RMS for the PU–AL condition. Likewise, we normalized upper limb EMG RMS amplitudes for each muscle (left AD, PD, BB, and TB; right AD, PD, BB, and TB) to intrasubject average EMG RMS for the AU–PL condition. We then averaged the intrasubject EMG RMS values to calculate the group EMG RMS value for each muscle per condition.

**Statistical Analysis**

For each question, we used a repeated-measure ANOVA (rmANOVA) to determine whether there were significant differences between conditions for each lower (or upper) limb muscle. For example, for question 1, we performed the rmANOVA looking at only the passive lower limb muscles for the PU–PL and AU–PL conditions. If the rmANOVA showed a significant difference among conditions, we used a Tukey Honestly Significant Differences (THSD) post hoc test to determine which condition(s) were significantly different (P < 0.05). For the single limb conditions, we used a chi-squared test of association (P < 0.05) to determine whether there was a significant difference in the number of muscles that had significant increases in EMG RMS with ipsilateral versus contralateral limb effort.

**RESULTS**

**Question 1: Does maximal effort in the upper (or lower) limbs increase muscle activation in passive lower (or upper) limbs?** Active upper limb effort resulted in increased muscle activation of the passive lower limbs. In a representative single-subject, the PU–PL condition had minimal muscle activation in the passive lower limbs, whereas the AU–PL condition had greater amplitudes and more distinct bursts in the passive lower limb muscle activation (Fig. 2A, black vs gray lines, respectively). When looking at the group average of all the subjects, the data showed the same effects as in the representative single-subject data (Fig. 2B). The AU–PL condition had significantly greater muscle activity than the PU–PL condition for the bilateral VM, MH, TA, and SO (Fig. 2C, asterisk over gray bars; THSD, P < 0.05). The knee joint angle data were similar for the two conditions, PU–PL and AU–PL (Fig. 2A and B). The handle and pedal forces indicated that subjects performed the stepping conditions as instructed, with the handle forces being substantially different and the pedal forces being similarly minimal. The small pushing pedal forces indicated the pedal pushing against the subject’s passive foot (Fig. 2A and B).

Similarly, lower limb muscle activation resulted in increased muscle activation of passive upper limb muscles. Representative single-subject data and group mean profiles showed burst-like muscle activity in the passive upper limbs during the PU–AL condition compared with the minimal muscle activity during the PU–PL condition (Fig. 3A and B, gray vs black lines). PU–AL EMG RMS amplitudes were significantly greater than PU–PL EMG RMS amplitudes for the bilateral PD, BB, and TB muscles (Fig. 3C, asterisk over gray bars; THSD, P < 0.05). The elbow joint angle, handle force, and pedal force data again indicated that subjects performed the stepping conditions as instructed (Fig. 3A and B) as the handle forces were similar although the pedal forces were different.
Question 2: Does simultaneous upper and lower limb maximal effort increase lower (or upper) limb muscle activation compared with only lower (or upper) limb maximal effort?

During simultaneous upper and lower limb maximal effort, there was no difference in muscle activation amplitudes compared with only upper limb maximal effort or only lower limb maximal effort. Representative single-subject data and group EMG mean profiles showed that muscle activation in the active lower limbs for the PU–AL and AU–AL conditions were similar in amplitude and shape (Fig. 4A and B, black vs gray, respectively). There were no significant differences in lower limb EMG RMS between the PU–AL and AU–AL conditions (Fig. 4C; rmANOVA, P > 0.05). Similarly,
there were no significant differences in upper limb EMG RMS between AU–PL and AU–AL (rmANOVA, \( P > 0.05 \)). The knee joint angle, handle force, and pedal force data indicated that subjects performed the stepping conditions as instructed (Fig. 4A and B).

**Question 3:** Does single limb maximal effort increase muscle activation in the ipsilateral or contralateral limb? Single upper limb active effort resulted in increased muscle activation in the passive ipsilateral lower limb compared with the contralateral lower limb. Representative single-subject data (Fig. 5, black) and group EMG mean profiles (Fig. 6A, black) for the Active Left Upper and Passive Lower condition showed clear rhythmic burst activity in the ipsilateral VM, ipsilateral TA, ipsilateral SO, and contralateral MH. When focused on just pulling during the Active Left Upper Pulling & Passive

![Figure 4](http://www.acsm-msse.org)
Lower condition, the EMG data had clear rhythmic burst activity in the ipsilateral VM, ipsilateral SO, and contralateral MH (Figs. 5 and 6A, light gray). When focused on just pushing during the Active Left Upper Pulling & Passive Lower condition, the EMG data had clear rhythmic burst activity in the ipsilateral TA (Figs. 5 and 6A, dark gray).

The knee joint angle, handle force, and pedal force data indicated that subjects performed the stepping conditions as instructed (Figs. 5 and 6A).

The EMG RMS amplitudes of the ipsilateral VM, MH, TA, and SO muscles were greater during the Active Left Upper & Passive Lower condition compared with PU–PL (Fig. 6B, asterisk in the Left Push & Pull column; THSD, \( P < 0.05 \)). The left MH and TA EMG RMS were also
significantly greater during the contralateral Active Right Upper & Passive Lower condition compared with the PU–PL amplitudes (Fig. 6B, asterisk in the Right Push & Pull column; THSD, $P < 0.05$). In the Pull Only instructions, Active Left Upper Pulling & Passive Lower resulted in significantly greater EMG RMS amplitudes in the ipsilateral VM and SO compared with PU–PL (Fig. 6B, asterisk in the Left Pull Only column; THSD, $P < 0.05$). The left MH had significantly greater EMG RMS during the contralateral Active Right Upper Pulling & Passive Lower condition.
DISCUSSION

We found three novel features of interlimb coupling on muscle activation during rhythmic movement in neurologically intact individuals in this study. Our first finding was a bidirectional coupling of muscle activation between upper and lower limbs. Maximum voluntary effort in the upper limbs increased passive muscle activation in the lower limbs. Likewise, maximum voluntary effort in the lower limbs increased passive muscle activation in the upper limbs. Our second finding was that interlimb neural coupling did not increase or decrease the total muscle recruitment possible during maximally activated lower (or upper) limbs. When subjects exerted simultaneous maximum upper and lower limb effort, there was neither facilitation nor inhibition of muscle activation compared with exclusively upper limb effort or exclusively lower limb effort. Our third finding was an ipsilateral coupling of muscle activation between upper and lower limbs. Maximum voluntary effort in a single upper limb increased muscle activation more in the ipsilateral lower limb than in the contralateral limb. This ipsilateral effect was also bidirectional. Single lower limb active effort also resulted in greater muscle activation in the ipsilateral passive upper limb (not shown). These results provide a more thorough understanding of the features and limitations of interlimb coupling of muscle activation between upper and lower limbs.

The general result that active effort in one limb pair resulted in increased muscle activation in passive muscles in the other limb pair was robust. It was consistent with our previous studies (15,17), held for single upper limb active effort conditions (Figs. 5 and 6), and was bidirectional (i.e., upper to lower, Fig. 2, and lower to upper, Fig. 3). This bidirectional effect agrees with studies examining the role of arm movement on cutaneous reflexes in the legs and the role of leg movement on reflexes in the arms (4,12,34). The increases in passive muscle activation with active effort in another limb(s) were small but significant. This agrees with an arm–leg cycling study that demonstrated a nontrivial subtle effect of arm cycling on lower limb reflexes even in the presence of the more dominant effect of leg cycling on lower limb reflexes (4).

The single limb conditions revealed a surprising ipsilateral, rather than contralateral, coupling of muscle activation. The ipsilateral coupling was evident in both directions, upper to lower (Figs. 5 and 6) and lower to upper. This ipsilateral coupling is interesting because one might suspect contralateral coupling between the upper and lower limbs. Reflex studies suggest that contralateral upper to lower limb coupling may be more prevalent during rhythmic movements compared with ipsilateral upper to lower limb coupling, although the relative strengths are uncertain in humans (31,33). The nervous system also naturally prefers in-phase movements (isodirectional) of ipsilateral limbs rather than antiphase movements (2,27). This suggests that there would be a preference for contralateral coupling during recumbent stepping because the contralateral upper limb moves in-phase with the lower limb, whereas the ipsilateral upper limb moves antiphase with the lower limb. Despite the antiphase movements of the ipsilateral limbs, we found a greater increase in muscle activation in the passive ipsilateral lower (or upper) limb during single active upper (or lower) limb effort (Figs. 5 and 6). The preference for ipsilateral coupling of wrist and ankle flexion has been attributed to coupled corticospinal drive rather than afferent signals associated with limb movements (1,5). On the basis of reflex studies, we might expect to see a preference for contralateral coupling if spinal mechanisms are dominant. Our results suggest that supraspinal drive may be more dominant compared with spinal mechanisms during a maximal effort rhythmic upper and lower limb task.

It is likely that a combination of neural mechanisms contribute to our observations of increases in passive muscle activation. One possible contributor is spinal connections between upper limb neural networks and lower limb neural networks (7,33). These spinal connections allow information about muscle activation in one limb to modulate
interlimb reflexes, coordination, and muscle activation of other limbs in both neurologically intact individuals and individuals with spinal cord injury (4,8,9,11–13,18,29). Another possible contributor is sensory feedback from one or more moving limbs, which modulates neural activity in another limb (6). Stretch reflexes, however, do not seem to contribute to our results because the timing and shape of the increased muscle activation during the passive limb conditions were similar to the timing and shape of the active limb conditions (15). Because recumbent stepping has been shown to have similar neural control to walking (26,32), locomotor commands from central pattern generators and/or supraspinal centers and the propriospinal pathways that modulate locomotor commands are also possible neural mechanisms for our findings. Another possible contributor is descending supraspinal drive that results in “cross-talk” and unintended muscle activation (2,10). Postural adjustments may also have a role in our observation of increased passive muscle activation; however, we stabilized the torso as much as possible using a torso strap. We previously tested torso stabilization with multiple straps but still observed increases in passive muscle activation. Regardless, anticipated postural demands required to stabilize the body during the maximal effort tasks cannot be excluded as a possible contributor. Future studies would need to use more extensive electrophysiological measurements to further delineate the possible pathways involved.

An important result of this study was that adding simultaneous maximal upper limb effort with maximal lower limb effort did not enhance lower limb muscle activation in neurologically intact subjects. The presumed spinal connections that contributed to increased passive muscle activation were not a significant factor when lower limbs were maximally activated. During the simultaneous upper and lower limb condition, most subjects were able to attain but not exceed the muscle activation levels of the only upper limbs or only lower limbs conditions (Fig. 4). The mean force results (not shown) paralleled our EMG RMS results showing that simultaneous upper and lower limb effort did not produce a significant decrease in handle or pedal forces compared with the conditions with only upper limb or only lower limb effort. This agrees with a previous study that showed no difference between unilateral and bilateral force production in a static task for an upper limb and contralateral lower limb pair (14). These results suggest that a phenomenon such as a bilateral deficit does not occur during a simultaneous maximum effort upper and lower limb rhythmic task of nonhomonymous muscles. Our active lower limb results (Fig. 4) also compliment an arm and leg cycling study that demonstrated no significant differences in lower limb EMG during a combined arm and leg cycling task to only leg cycling (4). This also suggests that humans are able to generate maximum voluntary muscle activation and force production during rhythmic tasks whether using both upper and lower limbs, only upper limbs, or only lower limbs.

A limitation of this study was the subject’s ability and motivation to perform the task as instructed. Because the device was always moving, there was no incentive to do any work during passive conditions. There was no reason to “cheat” and use the passive limbs to aid the active limbs. On the other hand, because the device was always moving, the only motivation for full effort was to comply with the instructions given. When subjects used maximal effort, the resistance increased to maintain the specified stepping frequency and the subject had to do more work. For our experiment, we only provided subjects with verbal encouragement to use maximal effort or stay relaxed during the data collections. We chose not to provide some form of biofeedback such as a display showing handle and pedal forces generated during the collection. Giving subjects biofeedback could allow subjects to voluntarily negate any underlying natural neural coupling between the upper and lower limbs through increased supraspinal activity.

Another limitation with this experiment was that we did not examine what happens at submaximal effort. There is still the possibility that upper limb exertion can increase muscle activation in lower limbs during submaximal exercise despite our results at maximal efforts. Transcranial magnetic stimulation could be used to determine whether there is an excitatory interlimb coupling during submaximal muscle activation that results in less supraspinal descending neural drive to the lower limbs with upper limb exertion (35). Understanding the role of upper limb effort on lower limb muscle activation at submaximal efforts has potential rehabilitation implications. This is of interest because therapeutic exercise and activities of daily living are often performed at submaximal levels. Perhaps incorporating active effort from unimpaired limbs may increase or improve muscle activation patterns in impaired limbs during whole body rhythmic tasks such as walking.

We conducted this study to answer questions about the effect of interlimb neural coupling on muscle activation. We found that maximum voluntary effort in the upper (or lower) limbs did increase muscle recruitment during passive rhythmic movement of the lower (or upper) limbs. Single limb effort conditions revealed a stronger ipsilateral coupling of muscle activation compared with contralateral coupling. We also found that simultaneous maximum upper and lower limb effort produced neither more nor less muscle activation compared with exclusively upper limb effort or exclusively lower limb effort in the active limbs. These results showed that active effort in one limb(s) can influence muscle activation in other unintended passive limbs during a rhythmic locomotor-like task in neurologically intact individuals. However, the presumed excitatory neural coupling did not enhance maximum activation. There are several factors that may contribute to this ipsilateral facilitatory neural coupling, including interlimb spinal neural connections that provide pathways for muscle activation from one limb(s) to contribute to muscle recruitment in another limb. These results support the existence of neural
connections between the upper and lower limbs and demonstrate that muscle activation in one limb(s) increases muscle activation in another limb(s) during a whole-body rhythmic task. Further studies examining the role of upper limb effort on lower limb muscle activation patterns will provide more insight into the potential of integrating upper limb effort during lower limb rehabilitation.

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