Features of hand-foot crawling behavior in human adults

M. J. MacLellan, Y. P. Ivanenko, G. Cappellini, F. Sylos Labini and F. Lacquaniti


You might find this additional info useful...
Features of hand-foot crawling behavior in human adults

M. J. MacLellan,1 Y. P. Ivanenko,1 G. Cappellini,1 F. Sylos Labini,1 and F. Lacquaniti1,2,3
1Laboratory of Neuromotor Physiology, Santa Lucia Foundation, Rome; 2Center of Space Bio-medicine and 3Department of Neuroscience, University of Rome Tor Vergata, Rome, Italy

Submitted 25 July 2011; accepted in final form 30 September 2011

ALTHOUGH BIPEDAL LOCOMOTION is a hallmark of humans, they can also walk on all four limbs. Toddlers often crawl before they begin walking unsupported; most of them crawl on the hands and knees, but some use a hands-and-feet pattern (Hildebrand 1967). Moreover, a few families have been identified with congenital central nervous system (CNS) disorders resulting in equilibrium impairment and a preferred locomotion on hands and feet (Getchell et al. 2001; Hildebrand 1967; Sparrow 1989; Sparrow and Newell 1994; Webb and Sparrow 2007). Even fewer studies have examined muscle activity during hands-and-knees crawling in a limited number of muscles (de Seze et al. 2008; Gallagher et al. 2011). Patrick et al. (2009) performed a thorough coordination analysis of crawling on hands and knees, as well as on hands and feet, in human adults and children. They found a large variability of limb coordination, suggesting flexible underlying control mechanisms. They concluded that humans may have an underlying neural circuitry similar to that of other primates and nonprimate quadrupeds, since many features of interlimb coordination are shared between these groups.

The coordination between the upper and lower limbs in humans may be due to sustained neural connections between the cervical and lumbosacral pattern generators (Wannier et al. 2001; Zehr et al. 2009). Such connections between the upper and lower limb neural controllers have been shown in humans by studying the modulation of electromyographic (EMG) and reflex activity of upper limbs muscles during rhythmic activity of lower limbs muscles and vice versa (Zehr et al. 2009), as well as by showing a tight coordination between the limbs in a variety of locomotor tasks (Dietz and Michel 2009; Donker et al. 2002; Ivanenko et al. 2005). Coupling between the upper and lower limbs has also received significant attention in the context of locomotor impairments (Crenna et al. 2008; Dietz 2002; Ferris et al. 2006; Klimstra et al. 2009; Tan 2010).

Taking the above considerations into account, we wondered whether reconstructing the spatiotemporal organization of the spinal motor output (by mapping muscle activations to the motoneuron pools) would provide insight into the coordination between the cervical (arm muscle control) and lumbosacral (leg muscle control) enlargements during quadrupedal and bipedal gaits in humans. We detailed the spatiotemporal characteristics of kinematic and muscle activation patterns when healthy adults crawled on their hands and feet at different speeds and at different support surface inclinations to generate a repertoire for comparison with human bipedal walking and quadrupedal walking in animals. The results show some unique features of quadrupedal gait performance in humans, including a flexible coupling between cervical and lumbosacral enlargement activity, differences between ground reaction force profiles between the upper and lower limbs, and the level of motoneuron activation in the spinal cord.

METHODS

Two different groups of participants volunteered for two separate experimental sessions. Ten healthy adults (7 males and 3 females, 32.6 ± 8.9 yr of age (mean ± SD), 74.6 ± 13.8 kg) were tested for the first session, and nine healthy adults (5 males and 4 females, 32.9 ± 8.3 yr of age, 67.6 ± 9.6 kg) were tested for the second session, with...
six participants common to both sessions. The studies were in accordance with the Declaration of Helsinki, and informed consent was obtained from all participants according to our procedures, which were approved by the Ethics Committee at the Santa Lucia Foundation.

Experimental Setup

The subjects participated in two experimental sessions: 1) treadmill locomotion at different speeds and inclinations and 2) overground locomotion at normal, slow, and fast speeds.

Session 1. Two experimental conditions were performed in this session. The first condition was level crawling or normal erect walking on a treadmill (EN-TRED; Enraf Nonius, Rotterdam, The Netherlands) at speeds ranging from 0.5 to 3.0 km/h in increments of 0.5 km/h (normal walking was also collected at 2, 3, and 5 km/h) with a rest period of ~3 min between trials. Participants were given time to adapt to each condition and provided a verbal signal to start collection. Each collection consisted of 10 consecutive strides. We also recorded crawling when the speed of the treadmill was ramped from 0.5 to 3 km/h in 25 s (ramp speed condition). The maximal speed (3 km/h) was chosen to correspond approximately to a comfortable and natural speed for crawling. The second condition consisted of crawling on the treadmill at 2 km/h with modest inclinations of 13°, 27°, and 35° (which avoided slippage of the feet on the treadmill surface). Again, participants provided a signal to start the trial, and 10 consecutive strides were collected. All participants performed the level crawling and the 13° inclination conditions, whereas 6 of 10 participants also performed the 27° and 35° inclinations. The total duration of the session was ~1.5 h. All participants were instructed to crawl on their hands and feet (as opposed to crawling on the hands and knees; see Gallagher et al. 2011; Patrick et al. 2009; Yokochi et al. 1990).

Session 2. Participants crawled overground at self-paced normal, slow, and fast speeds while ground reaction forces and kinematics data were recorded. The crawling direction was oriented such that only ipsilateral (right side) ground reaction forces were recorded (Fig. 1B). For these trials, participants crawled at self-selected normal, slow, and fast speeds for 3 trials each or until there complete ground contacts were collected with the right forelimbs and hindlimbs.

Data Recording

Kinematics data were recorded with a 9-camera Vicon-612 system (Oxford, UK) at 100 Hz. Infrared reflective markers (diameter 1.4 cm) were placed bilaterally on the following anatomic landmarks: first distal phalanx, fifth metatarsal head, heel, lateral malleolus, lateral femoral epicondyle, greater trochanter, acromioclavicular joint, lateral humeral epicondyle, ulnar process, and distal phalanx of middle finger (Fig. 1A). All marker data were low-pass filtered offline using a second-order dual-pass Butterworth filter and bandstop filtered at a frequency of 50 Hz. EMG data were then full-wave rectified and low-pass filtered using a second-order dual-pass (zero phase) Butterworth filter with a cutoff of 10 Hz.

Ground reaction forces were collected (session 2) with a force plate (0.9 × 0.6-m 9287B; Kistler, Zurich, Switzerland) at 1,000 Hz. All ground reaction force data were low-pass filtered offline using a second-order dual-pass Butterworth filter with a cutoff of 20 Hz, normalized to participant body weight, and time-interpolated to 100 points for interstride averaging.

Data Analysis

The recorded kinematics data were used to construct a linked segment model consisting of the trunk, upper arm, forearm, hand, thigh, shank, and foot bilaterally. This model was used to calculate joint angles on the right side of the body at the wrist, elbow, and shoulder in the upper limb and the ankle, knee, and hip in the lower limb (Fig. 1A, arrows indicate positive joint motion). Cycle durations of the upper and lower limbs were determined by two consecutive right wrist contacts and first distal phalanx contacts, respectively (the first parts of the body to make contact with the ground during crawling). Swing duration of the upper and lower limbs was determined from lift off of the right distal phalanx of middle finger to wrist contact and lift off of the right first distal phalanx to first distal phalanx contact, respectively. Excursions of the upper and lower limbs were determined by the distance traveled by the right wrist and first distal phalanx, respectively, during swing. Ipsilateral phase lag (IPL) between upper and lower limbs was determined using the methods described by Patrick et al. (2009). In brief, the relative timing of right upper limb contact was expressed as a percentage of the gait cycle determined by consecutive right foot contacts:

$$\text{IPL} = \frac{t_1}{T} \times 100\%$$

where $t_1$ is interval of time between right foot and right hand touchdown events and $T$ is cycle duration (Fig. 2A). According to this method, lateral gait patterns (ipsilateral upper/lower limb contact at similar instances) are determined at a value of 0% and diagonal gait patterns (contralateral upper/lower limb contact at similar instances) are determined at a value of 50%. Intermediate values (~25%) correspond to no limb pairing.

The orientation of the upper limb (shoulder to point of contact, acromioclavicular joint-distal phalanx of middle finger) relative to a
vertical line through the proximal joint center at both touchdown and liftoff of a step were computed. Similarly, this was done for the lower limb (hip to contact point: greater trochanter-fifth metatarsal head). Vertical displacement of the hip (greater trochanter) and shoulder (acromioclavicular) joints were calculated as the vertical range over a stride cycle. Average limb orientation was determined for each step by calculating the difference between the limb angle at touchdown and at liftoff and then dividing this number by two. This midpoint of the total limb excursion defines the axis about which the limb oscillates during support phase, as well as allowing comparison with the same angle in selected animals (Larson and Stern 2009). An average limb angle greater than zero indicates a protracted limb axis, whereas values less than zero indicate a retracted limb axis.

Ground reaction force profiles were obtained from the right hand and foot and normalized to participant body weight. The axes of the force plate were aligned such that positive ground reaction forces were directed in the upward (vertical) and forward (horizontal) directions. For graphical purposes, ground reaction force data were time normalized and averaged across participants. Peak force was determined from the maximum (and minimum for horizontal forces) and five cervical (C5–T1) discrete activation waveforms to examine the output from the main enlargements of the spinal cord. The Kendall chart indicates only the segments innervating each muscle and not the proportion of total muscle activation attributable to each segment. For comparison with previously published cat data that include a greater spatial resolution (Yakovenko et al. 2002), we also used the Sharrard (1964) data table for innervation, which approximates the proportion of total muscle activation attributable to each segment (by taking multiple slices within each spinal segment), instead of assuming equal proportions in all segments. To this end, the contribution of each muscle to the estimated activity of the jth segment \( (Eq. 2) \) was multiplied by its PCSA (Cappellini et al. 2011).

We used the Kendall chart results in the six lumbosacral (L2–S2) and five cervical (C5–T1) discrete activation waveforms to examine the output from the main enlargements of the spinal cord. The Kendall chart indicates only the segments innervating each muscle and not the fraction of total motor pool of the muscle that can be assigned to a segment. For comparison with previously published cat data that include a greater spatial resolution (Yakovenko et al. 2002), we also used the Sharrard (1964) data table for innervation, which approximates the proportion of total muscle activation attributable to each segment (by taking multiple slices within each spinal segment), instead of assuming equal proportions in all segments. To this end, we subdivided each segment into six subsegments and applied the same equation \( (Eq. 2) \) for each jth subsegment (Ivanenko et al. 2006). The resulting spinal cord maps of activation were not smoothed but contained 36 discrete bands (6 subsegments × 6 segments; L2–S2).

The recorded patterns of EMG activity were mapped onto the approximate rostrocaudal location of motoneuron (MN) pools in the human spinal cord (Ivanenko et al. 2006). This approach provides an interpretation of the output from locomotor central pattern generators (CPGs) in terms of segmental control rather than in terms of individual muscle control (Yakovenko et al. 2002). This approach can be used to characterize network architecture for different gaits by considering relative intensities, spatial extent, and temporal structure of the spinal motor output (Ivanenko et al. 2008).

In this study, we used two myotomal charts: those of Kendall et al. (1993) and those of Sharrard (1964). Kendall et al. compiled reference segmental charts for all body muscles by combining the anatomic and clinical data from six different sources. A capital X in Kendall’s chart denotes localization agreed on by five or more sources, a lowercase x denotes agreement of three to four sources, and a bracketed (x) denotes agreement of only two sources. In our maps, X and x were weighted 1 and 0.5, respectively, whereas we discarded (x). We assumed that our population of subjects had the same spinal topography as this reference population.

To reconstruct the output pattern of any given spinal segment \( S_j \) of the most active lumbosacral segments (L2–S2), we averaged all rectified EMG waveforms corresponding to that segment (Cappellini et al. 2010; Ivanenko et al. 2008):

\[
S_j = \frac{\sum_{i=1}^{n_j} k_{ij} \cdot EMG_i}{n_j}
\]

where \( n_j \) is the number of EMG waveforms corresponding to the jth segment, \( k_{ij} \) is the weighting coefficient for the ith muscle (X and x in Kendall’s chart were weighted with \( k_{ij} = 1 \) and \( k_{ij} = 0.5 \), respectively). The assumption implicit in this method is that the rectified EMG provides an indirect measure of the net firing of MNs of that muscle. EMGs were expressed in microvolts and normalized to the physiological cross-sectional area (PCSA) of lower limb (Ward et al. 2009) and upper limb muscles (Ramsay et al. 2009; Wood et al. 1989). To this end, the contribution of each muscle to the estimated activity of the jth segment \( (Eq. 2) \) was multiplied by its PCSA (Cappellini et al. 2011).
To compare neural activation with crawling kinematics, we performed a cross-correlation between the dominant MN activity in the lumbar (sum of activity from L2 and L3) and cervical enlargements (sum of activity from C8 and T1). To determine the phase shift between lumbar and cervical activity, we constructed an algorithm to find the largest correlation coefficient (which was generally above 0.8) and expressed the phase shift as a percentage of the gait cycle.

Statistics

Descriptive statistics include means ± SD. Repeated-measures ANOVA was used to assess the effect of speed and crawling conditions on gait and motor pattern parameters as well as differences between upper and lower limbs. Paired t-tests were used to determine differences between upper and lower limbs for peak forces and impulses, and a one-tailed t-test was used to determine whether average limb angle differed from zero. A Pearson correlation was used to test statistically the relationship between lumbar-cervical activity phase shift and IPL. Reported results are considered significant for \( P < 0.05 \).

RESULTS

Basic Gait Parameters

Figure 2A illustrates a typical footfall pattern in human crawling. In general, four, three, or two limbs were simultaneously supporting body weight depending on the speed and the instant of the gait cycle. Subjects tended to use a lateral footfall sequence when crawling over the studied range of speeds, i.e., the footfall of a lower limb usually followed that of the ipsilateral upper limb. We quantified this coordination pattern (Fig. 2B) by computing the contact of the right upper limb expressed as a percentage of the gait cycle (Patrick et al. 2009). The results of the IPL analysis between the upper and lower limbs did not differ significantly as a function of crawling speed \( [F_{(5,45)} = 0.204, P = 0.87, \text{Fig. 2B, left}] \) or inclination \( [F_{(3,12)} = 1.406, P = 0.30, \text{Fig. 2B, right}] \), although a range of different IPL values was observed across participants. Overall, participants showed no limb pairing (IPL ~25%). However, there were participants who exhibited a more “diagonal” (IPL close to 50%) or “lateral” pattern (IPL close to 0%), and this pattern did not change between crawling conditions (Fig. 2).

As expected, cycle duration decreased \( [F_{(5,45)} = 113.558, P < 0.001] \) and percentage of swing time increased significantly \( [F_{(5,45)} = 109.210, P < 0.001] \) with changes in treadmill speed (Fig. 3A). Although cycle duration did not differ significantly between the upper and lower limbs \( [F_{(1,9)} = 0.874, P = 0.37] \), swing duration was consistently shorter in the upper compared with the lower limbs \( [F_{(1,9)} = 13.96, P = 0.005] \). Moreover, limb excursion increased significantly with treadmill speed \( [F_{(5,45)} = 10.875, P = 0.002] \) and was systematically larger in the upper limb \( [F_{(1,9)} = 8.897, P = 0.015] \), consistent with longer stance durations for the arms.

Changing the inclination of the crawling surface had no significant effect on cycle duration \( [F_{(3,15)} = 0.174, P = 0.77] \), percentage of swing duration \( [F_{(3,15)} = 0.359, P = 0.58] \), or limb excursion \( [F_{(3,15)} = 0.236, P = 0.70] \), and differences were not evident between upper and lower limbs for cycle \( [F_{(1,5)} = 1.007, P = 0.36] \), swing duration \( [F_{(1,5)} = 0.317, P = 0.64] \), or limb excursion \( [F_{(1,5)} = 0.084, P = 0.78] \) (Fig. 3B).

Figure 4 illustrates kinematic patterns in human crawling on hands and feet. Vertical greater trochanter (hip) displacements were significantly higher \( [F_{(1,7)} = 34.192, P < 0.001] \) than those of the acromioclavicular joint (shoulder) at all speeds, although no significant differences were observed between treadmill speeds \( [F_{(5,35)} = 0.554, P = 0.73] \) (Fig. 4, A and B, top). Among the joint angles, the shoulder \( [F_{(5,35)} = 40.035, P < 0.001] \) and elbow \( [F_{(5,35)} = 4.939, P = 0.021] \) joint angle ranges increased with speed, whereas speed effects were not evident at all other joints \( (P > 0.05) \) (Fig. 4, A and B, bottom).

The orientation of the lower limb differed greatly from that of the upper limb (Table 1). The lower limb was greatly retracted for all crawling speeds \( [F_{(1,8)} = 70.221, P < 0.001] \). In contrast, the upper limb became more retracted as crawling speed increased \( [F_{(5,40)} = 29.590, P < 0.001] \) with a 15 ± 2° protraction at 0.5 km/h and a 9 ± 2° protraction at 3 km/h, whereas no significant changes were found for the retraction of the lower limb \( [F_{(5,45)} = 0.474, P = 0.64] \).

Human crawling on feet is characterized by distinct patterns of propulsive forces for the lower limbs and braking forces for the upper limbs (Fig. 5A). Comparisons of peak force indicated greater vertical force in the lower compared with the upper limb \( (P < 0.001) \) and significantly greater peak positive and negative forces for the lower and upper limbs, respectively \( (P < 0.001 \text{ in both cases}) \) (Fig. 5B). The propulsive and braking function of the legs and arms, respectively, became even more obvious if positive and negative impulses of shear forces were compared: the positive thrust for the legs was on average 10 times greater than the negative one; for the arms, the difference was even larger (15 times). Over a wide range of crawling speeds (Fig. 5C), one can see that peak vertical and horizontal forces tend to increase with crawling speed in the lower
EMG Activity and MN Activity

Figure 6A illustrates ensemble-averaged EMG waveforms through the stride cycle for crawling at increasing speeds. The largest differences in EMG activity relative to erect walking were in the upper limbs where there is a large burst of activity in the deltoids and triceps during upper limb stance. At the wrist, we saw that extensor activity increased during upper limb swing, whereas flexors exhibited a single burst at the end of upper limb stance. In the lower limbs, there was a trend for quadriceps (rectus femoris, vastus lateralis, vastus medialis) and tibialis anterior EMG activity to increase with crawling speed, whereas the burst in the ankle plantar flexors (lateral gastrocnemius, medial gastrocnemius, soleus) was diminished compared with upright walking.

With reference to cervical MN activity patterns in Fig. 6A, the activity was focused around the C8 and T1 segments during upper limb support with minimal activation during swing. The wrist flexor burst at the end of upper limb stance is shown by activity in the C6–T1 segments. In the lumbosacral region (Fig. 6A, bottom), activity was focused around the L2–L4 segments immediately before and during lower limb stance. During swing, there was also a burst of activity around L4 and L5 that tended to increase with crawling speed. The average activations over the stride cycle for the cervical and lumbosacral enlargements at each crawling speed are shown in Fig. 6B. Statistical analysis showed that MN activity increased significantly with crawling speed \( F_{(5,45)} = 31.515, P < 0.001 \) but did not differ significantly between cervical and lumbosacral enlargements \( F_{(1,9)} = 0.268, P = 0.62 \).

In general, the total MN activation level (and muscle activity) was considerably higher in crawling than in erect walking for both the cervical and lumbosacral enlargements (Fig. 6B). It is also worth noting that the participants verbally stated that the crawling protocol was much more fatiguing than normal walking.

Although temporal MN activation differed somewhat between participants (typical participants who crawled with lateral, no limb pairing, or diagonal patterns are shown in Fig. 7A), this was related to crawling behavior. With comparison of dominant MN patterns with crawling kinematics, a strong relationship \( r^2 = 0.866, P < 0.001 \) was shown between the cervical-lumbar spinal segment activity phase difference and IPL (Fig. 7B). It should be noted that two data points from a single participant were removed due to low levels of activation in the lumbar spinal segments.

Effect of Changes in the Surface Inclination

With increasing inclination of the crawling surface, we observed further changes in the EMG activity patterns (Fig. 8A). During upper limb stance, there was a decrease in triceps and posterior deltoid activity and even greater decreases in anterior deltoid activity as surface inclination increased. Although we observed these changes in shoulder musculature, the activation of wrist extensors during upper limb swing and the burst of wrist flexor activity at the end of upper limb stance remained in all conditions. In the lower limb, the quadriceps and tibialis anterior activity was similar at all inclinations, but there was a gradual appearance of gastrocnemius and soleus activity as inclination increased.

The changes in upper limb muscle activity are shown in terms of different MN burst patterns as surface inclination increased (Fig. 8A). In particular, cervical activity throughout

**Table 1. Average limb orientation in mammals during quadrupedal walking**

<table>
<thead>
<tr>
<th>Group</th>
<th>Average FL. Angle, deg</th>
<th>Average HL. Angle, deg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small mammals</td>
<td>-6 ± 5 (6)</td>
<td>3 ± 7 (6)</td>
</tr>
<tr>
<td>Primates</td>
<td>-2 ± 3 (50)</td>
<td>2 ± 4 (47)</td>
</tr>
<tr>
<td>Humans</td>
<td>9 ± 2* (10)</td>
<td>-21 ± 2† (10)</td>
</tr>
</tbody>
</table>

Values are means ± SD; numbers in parentheses are the number of subjects. FL, forelimb; HL, hindlimb. Positive values denote limb protraction, zero refers to the vertical average limb orientation. The data for small mammals and primates are Larson and Stern (2009). The data on human crawling are presented at 3 km/h. A 1-tailed t-test was applied to assess significant differences from zero. *\( P < 0.0001 \); †\( P < 0.000001 \).
upper limb stance in the C8 and T1 segments diminished, with a burst of activation in the C6–T1 segments at the end of upper limb stance persisting with increasing surface inclination. In the lumbosacral region, activity in the L3 and L4 segments remained and there was an appearance of a burst in the S1 and S2 segments at the end of lower limb stance that increased with surface inclination (Fig. 8A, bottom). The average motoneuron activity throughout the stride is illustrated in Fig. 8B, and statistical analysis showed a surface inclination by spinal segment interaction effect \[F(6,30) = 15.784, P < 0.001\], suggesting an increase of sacral activity and a decrease in cervical activity as surface inclination increases.

**DISCUSSION**

This study specifically examined the spatiotemporal pattern of muscle activity, general spinal motor output characteristics, and kinematic and kinetic determinants of human crawling on hands and feet. Our results showed a number of similarities with quadrupedal locomotion in animals, but they also revealed several features specific to humans. Below we discuss our findings in the context of neuromuscular control of the lower and upper limbs and its comparison with animal locomotion.

**Motor Patterns in Human Crawling**

Specialized neural circuits in the caudal and rostral spinal cord organize the hindlimb and forelimb locomotor activity during animal locomotion (Grillner 2006; Kiehn 2006; Zehr et al. 2009). The coordination between cervical and lumbosacral pattern generation circuitry may be provided via mechanical coupling of arm-trunk-leg movements (Falgairolle et al. 2006; Pearson 2001) and via the long projecting propriospinal neurons coupling the cervical and lumbar spinal cord (Nathan et al. 1996). In this study we used an approach that provides information about CPG output in terms of segmental control rather than in terms of individual muscle control (Grasso et al. 2004; Ivanenko et al. 2006; Monaco et al. 2010; Yakovenko et al. 2002). Our results suggest that the temporal linkage between cervical and lumbosacral generators is flexible and gait dependent (Figs. 6–8). In this context, it is also noteworthy that cervical MN activity exhibits different timings during human
Fig. 6. Motor patterns in human crawling at different speeds. A: ensemble-averaged patterns of electromyographic (EMG) activity across participants (n = 10). Mean waveforms are plotted against the normalized gait cycle (beginning from the foot contact of the right leg). Corresponding α-motoneuron (α-MN) activation patterns (normalized to physiological cross-sectional area method using Kendall’s chart) of cervical and lumbosacral spinal enlargements are shown at the bottom of each panel. Bic, long head of biceps brachii; Tri, long head of triceps brachii; DeltA, anterior deltoit; DeltP, posterior deltoit; LD, latissimus dorsi; Pec, pectoralis major; BR, brachioradialis; ECU, extensor carpi ulnaris; ECR, extensor carpi radialis; FCU, flexor carpi ulnaris; FCR, flexor carpi radialis; FDS, flexor digitorium superficialis; GM, gluteus maximus; Gmed, gluteus medius; TFL, tensor fascia latte; Sart, sartorius; AddL, adductor longus; RF, rectus femoris; VL, vastus lateralis; VM, vastus medialis; ST, semitendinosis; BF, biceps femoris; LG, lateral gastrocnemius; MG, medial gastrocnemius; Sol, soleus; TA, tibialis anterior. B: group means (±SD) of lumbosacral (left) and cervical (right) activations from crawling and walking.
walking vs. running (Ivanenko et al. 2008), likely due to differences in arm configuration and movements in these two gaits (flexed elbows during running). The coupling of the cervical and lumbosacral enlargements and the relative stance/swing durations are gait and context dependent (Wannier et al. 2001), consistent with the neuromechanical tuning hypothesis according to which the phase durations generated by the CPGs closely match the required kinematics (Prochazka and Yakovenko 2007; Sylos Labini et al. 2011). Compared with habitual quadrupeds, human crawling may demand more cortical control due to the unfamiliar body posture requirements and/or intersegmental coordination. Furthermore, crawling experience may also contribute to the cortical control (Bell and Fox 1996).

The data in the literature on human crawling are rather limited. Webb et al. (2007) using harmonic analysis suggested that the arms behave like a pendulum in bipedal, but not quadrupedal, human locomotion. Swing phase in the upper limbs is to absorb energy to decelerate the body, whereas the lower limbs generate energy to progress the body forwards (Hanna et al. 2006). Nonhuman primates also show a more important propulsive role of the hindlimbs compared with the forelimbs (Kimura 1985), although to a much lesser extent than humans do (Fig. 5). Moreover, in human crawling, the lower limbs are retracted more than the hindlimbs in small mammals and primates (Table 1), presumably reflecting a relatively longer lower limb length and/or specific limb segment proportions in humans (Ivanenko et al. 2011; Leurs et al. 2011; Polk 2004).

Finally, a remarkable feature of human crawling is a lack of ankle plantar flexor muscle activation and, as a consequence, a relatively lower activation of sacral segment motoneuron output. The double-humped pattern of vertical ground reaction force seen in cats (Lavoie et al. 1995) and in human erect walking (Winter 1991) is not observed in human crawling. The horizontal ground reaction force reveals predominantly negative and positive patterns in the upper and lower limbs, respectively (Fig. 5), consistent with the notion that the primary function of the upper limbs is to absorb energy to decelerate the body, whereas the lower limbs generate energy to progress the body forwards (Hanna et al. 2006). Nonhuman primates also show a more important propulsive role of the hindlimbs compared with the forelimbs (Kimura 1985), although to a much lesser extent than humans do (Fig. 5). Moreover, in human crawling, the lower limbs are retracted more than the hindlimbs in small mammals and primates (Table 1), presumably reflecting a relatively longer lower limb length and/or specific limb segment proportions in humans (Ivanenko et al. 2011; Leurs et al. 2011; Polk 2004).

**Comparison With Cat and Monkey Locomotion**

Our findings revealed both similarities and differences between the human patterns and those of cat and monkey locomotion. Among the similarities, one may point out the alternating activity of cervical and lumbosacral enlargements, which reflects a diagonal nature of the interlimb coordination in primate locomotion (such as that observed in rhesus monkeys by Courtine et al. 2005), although humans exhibit great variability in interlimb phasing (Fig. 2). The relative swing duration and excursion also increased with speed (Fig. 3) as they do in rhesus monkeys (Courtine et al. 2005). Moreover, activation patterns in muscles such as vastus lateralis, rectus femoris, gluteus medius, sartorius, and tensor fascia lata displayed prolonged bursts during stance (Fig. 6A) relative to bipedal walking, but this prolonged pattern is similar to that in cats (Yakovenko et al. 2002) and monkeys (Courtine et al. 2005).

Although human crawling shares a number of common gait features with these animals, we found several critical differences in ground reaction forces and limb orientation and a reorganization of spinal segment motoneuron output. The double-humped pattern of vertical ground reaction force seen in cats (Lavoie et al. 1995) and in human erect walking (Winter 1991) is not observed in human crawling. The horizontal ground reaction force reveals predominantly negative and positive patterns in the upper and lower limbs, respectively (Fig. 5), consistent with the notion that the primary function of the upper limbs is to absorb energy to decelerate the body, whereas the lower limbs generate energy to progress the body forwards (Hanna et al. 2006). Nonhuman primates also show a more important propulsive role of the hindlimbs compared with the forelimbs (Kimura 1985), although to a much lesser extent than humans do (Fig. 5). Moreover, in human crawling, the lower limbs are retracted more than the hindlimbs in small mammals and primates (Table 1), presumably reflecting a relatively longer lower limb length and/or specific limb segment proportions in humans (Ivanenko et al. 2011; Leurs et al. 2011; Polk 2004).
Fig. 8. Motor patterns in human crawling at different surface inclinations and during level upright walking at 5 km/h. The format of the presentation in A is similar to that of Fig. 6 with a separate representation of lumbar and sacral activity in B. The dashed lines in B represent mean lumbosacral and cervical α-MN activation during upright walking at 5 km/h.
segment proportions, muscle attachments, passive elasticity) is adapted to bipedal walking. Given the retracted limb orientation (Table 1), the ground reaction force vector may be closer to the ankle joint center, resulting in a decreased moment of force in the ankle joint. Nevertheless, a prominent activation of thigh muscles and a dominant activity of lumbar segments were present in human crawling at all speeds (Fig. 6). On the contrary, the available data in cats (Yakovenko et al. 2002) and monkeys (Courtine et al. 2005) indicate a more uniform redistribution of proximal and distal extensor activities and a prominent activation of the caudal spinal segments (Fig. 9, left). It should be stressed that the quadrupedal mode of walking we investigated in this study is, of course, different from that exhibited by gorillas and chimpanzees, which walk on their knuckles, and from orangutans, which walk on their fists. In addition, several research groups have trained monkeys to walk with a bipedal gait pattern (Fitzsimmons et al. 2009; Hirasaki et al. 2004; Nakajima et al. 2004). From the energetic point of view, nonhuman primates trained to walk bipedally consume less metabolic energy during quadrupedal than bipedal gait (Nakatsukasa et al. 2004), whereas humans display considerably higher levels of mean MN activation in both cervical and lumbosacral enlargements during crawling compared with walking (Fig. 6B), supporting the idea of a high specialization of the neuromuscular apparatus and body proportions to specific gaits (Alexander 2004; Ivanenko et al. 2011; Leurs et al. 2011; Polk 2004; Sockol et al. 2007).

**Crawling at Different Inclines**

We assessed motor patterns for three levels of upslope crawling and compared them with similar data for level treadmill crawling and walking (Fig. 8). The principal purpose of this upslope crawling paradigm was to gain insight about the functions of muscles and the mechanisms of neural control required for different body orientations and support surfaces. Also, crawling at increasing inclines represents a gradual transition toward the upright body orientation and thus an “intermediate” form of locomotion between quadrupedal and erect bipedal walking. Although these similarities exist, fundamental differences are also evident. As surface inclination increases, there is a decrease in cervical α-MN activity that leads to an overall decrement of upper limb muscle activity (Fig. 8B, right). More importantly, we see the appearance of a sacral α-MN burst and increased ankle plantar flexor muscle activation (Fig. 8, left), likely related to the change in direction of the ground reaction force relative to the ankle joint (due to the more horizontal foot segment orientation compared with level crawling, see above), leading to a larger moment of force. The reason for these changes may be directly related to the redistribution of body weight between the upper and lower limbs and across lower limb joints.

**CONCLUSIONS**

Overall, this study provides new data on the reorganization and adaptation of the output of central pattern generators.
involved in the control of quadrupedal human locomotion as a function of different environmental constraints. Of particular interest was a decrease in sacral compared with lumbar motoneuron activation and the flexibility in the spatiotemporal output for the spinal segments. Sacral activity tended to increase with increasing surface inclination, more closely resembling upright walking. The findings of neuromechanical coupling between upper and lower limb motion could lead to a better understanding of the hypothetical evolution of human gait from quadrupedal locomotion (Schmitt 2003; Thorpe et al. 2007; Zampagni et al. 2011) and also may be of particular interest in the construction of biologically inspired robots (Ijspeert 2008; Pfeifer et al. 2007) or for clinical applications (Ferris et al. 2006; Klimstra et al. 2009; Tan 2010).

**REFERENCES**


