Qualitative Comparison between Rats and Humans in Quadrupedal and Bipedal Locomotion

Taisei Hosoido¹, Futoshi Mori²*, Keita Kiyoto², Takashi Takagi², Yukari Sano², Megumi Goto², Katsumi Nakajima³, Naomi Wada²

¹Osaka Veterinary Referral Center, Osaka, Japan
²Laboratory of Veterinary System Science, Yamaguchi University, Yamaguchi, Japan
³Department of Physiology, Kinki University School of Medicine, Osaka-Sayama, Japan
Email: *morif@yamaguchi-u.ac.jp

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ABSTRACT

Bipedal (Bp) locomotion is one of the most characteristic motor behaviors in human beings. Innate quadrupedal (Qp) four-legged animals also often walk bipedally. The walking posture, however, is significantly different between the two. This suggests that although both have a potential to walk bipedally, however, the human has a body scheme suitable for Bp locomotion, probably its skeletal system. The skeletal system includes the lumbar lordosis, sacral kyphosis, a round pelvis, a large femur neck angle, short feet, and so on. To verify this hypothesis, we compared kinematic and EMG activities between rats and humans during Qp and Bp locomotion on a treadmill belt. The rat is a representative Qp animal, but it is able to acquire Bp walking capability with motor learning. Although the mobile ranges of the hindlimb joint are different during each locomotor pattern between rats and humans, both showed replicable flexion and extension excursion patterns for each joint depending on the locomotor phase. There are many phase-locked EMG bursts between rats and humans during the same walking task and these are observed in the proximal rather than the distal muscles. This suggests that both rats and humans utilize similar neuronal systems for the elaboration of Qp and Bp locomotion. It was interesting that both subjects showed more muscle activities during non-natural locomotor patterns; Qp < Bp for rats and Bp < Qp for humans. This indicates that rat Bp and human Qp walking need more effort and we may be able to find its reason in their skeletal system.

Keywords: Human; Rat; Evolution; Posture; Locomotion; Bipedal; Quadrupedal

1. Introduction

Most terrestrial vertebrates walk with the limbs on the ground and terrestrial locomotion can be divided into quadrupedal (Qp) and bipedal (Bp) locomotion. The majority of mammals walk quadrupedally, while humans walk bipedally. As Aristotle and Darwin showed, the upright standing posture and Bp walking are the most basic and essential features of human beings. For this, we have developed a unique skeletal system including the lumbar lordosis, sacral kyphosis, a round pelvis, large femur neck angle; short feet, and so on. The lumbar lordosis in humans is a critical component in the ability to achieve upright and bipedal locomotion [1]. It is shown that such skeletal adaptation suitable for human like Bp walking was evolved relatively recent [2]. As a result, humans are a unique mammal that can adapt their Bp walking patterns to the various walking circumstances. However, many other innate Qp four-legged animals (Qp animals) can also stand upright and walk bipedally without the skeletal characteristics observed in humans. Bears and even cats and dogs can stand on their hindlimbs and walk bipedally if trained for a circus attraction. It is also well known that some species of genetically Qp nonhuman primates walk bipedally in the wild. In this sense, the Bp locomotion itself is by no means limited to humans or some nonhuman primates.

What are the triggers for bipedalism? During Qp walking, the forelimbs are needed. Therefore, Qp animals support their body weight, maintain body balance and produce propulsive force with their limbs. However, Qp animals often use their forelimbs not only for the execution of locomotion, but also for other purposes. These include gripping and collecting food, hanging from branches, digging up soil to find food and nesting. These functional usages of the forelimb may have already existed when mammals appeared on the earth (Jurassic period) and reinforced the different functional roles between forelimbs and hindlimbs. The functional differences are often

*Corresponding author.
observed in the morphology. In this sense, one can easily observe skeletal differences between forelimbs and hindlimbs in the most primitive mammals, *Monotremes*, on earth. These are also observed in other relatively old mammals such as the Insectivora and Marsupials. For further development of such specific forelimb function, the forelimbs must be freed from locomotion by standing up.

It has been considered that human beings evolved from Qp animals. Because locomotor control mainly consists of skeletomuscular and neuronal systems in both Qp mammals and Bp human, the evolutionary transition from Qp to Bp walking may be primarily found in the changes in these systems [3]. Human infants start walking quadrupedally and then acquire stable Bp capability as they grow up. One possible reason why the walking pattern of human infants smoothly transfers from Qp to Bp walking is that the infants already have a skeletomuscular system suitable for Bp walking at birth. It is interesting that infants develop lordosis around the time that they acquire bipedal locomotion. Furthermore, the facts described above suggest that reorganization of the central nervous system (CNS) for bipedalism is not essentially required at birth in human beings. Moreover, nonhuman primates including chimpanzees cannot fully extend their hip and knee joints when they walk bipedally, and have a “bent-hip and bent-knee” walking posture [4]. This is because the skeletal structure around their ischia is not adequate for full extension of the hip joint. These observations strongly suggest that innately Qp mammals do not have a skeletal system suitable for the elaboration of Bp walking, but use an inherent neuronal system similarly for both Qp and Bp locomotion. However, the defining factor in the shift from Qp walking to Bp walking remains to be elucidated. Here, we test the hypothesis that four legged mammals and humans have their similar inherent neuronal systems for the elaboration of both Qp and Bp walking. To do this, we compared their kinematic characteristics and recorded electromyographic activities from different types of muscles during Qp and Bp walking. Rats are originally Qp animals, but we had showed that they are able to acquire Bp walking capability by training [5].

2. Materials and Methods

2.1. Subjects

Twenty-five rats, of both sexes were used and 5 healthy male, human subjects with no history of motility disorders volunteered for this study. We established rat Bp walking model (RBWM) to study the neuronal control mechanisms of Bp walking and the effects of Bp walking on the central nervous system [5,6]. One of the characteristic features in our model is that the forelimbs, which are often amputated when establishing a Bp animal model, were preserved. The detailed bipedal training method was described elsewhere [5]. Briefly, 3 weeks after birth, the rats were housed in a large rectangular glass container that required them to use an upright posture to reach water for 2 to 3 weeks. Water was provided through drip bottles that were placed on the top of the container. Then, they began training using a bipedal-walking training device. This device moves the drip bottle at a constant speed, and the rat follows the bottle to obtain water while maintaining an upright posture. Each rat underwent training for 30 mins/day and 7 days/week for at least 3 months. Water was given to the rats from the drip bottle installed on the bipedal-walking training device or treadmill during the training periods. All procedures confirmed to the guidelines of the Animal and Human Investigation Committee and the Ethics Committee at Yamaguchi University. Informed, written consent was obtained from each human subject. The experiments were performed according to the Declaration of Helsinki.

2.2. Behavioral Task and Data Analysis

2.2.1. Qp and Bp Walking Tasks

The RBWM and humans participated in the following two walking tasks.

**Qp walking:** Both RBWM and human subjects walked on the treadmill belts quadrupedally at a comfortable speed for at least one minute. Each collection consisted of 5 - 10 stable consecutive strides as a minimum number. For humans, all subjects were instructed to walk on their hands and feet and were given time to adapt to this Qp walking pattern on the treadmill belt before recording.

**Bp walking:** Both subjects walked on a treadmill belt bipedally at a comfortable speed for at least a half minute. In RBWM, each collection consisted of at least 3 - 5 stable consecutive strides as a minimum number. For human subjects, all subjects were given time to adapt their Bp walking on the treadmill belt before recording.

2.2.2. Data Recording

The right side lateral views of the Qp and Bp walking in both RBWM and humans were recorded using a high speed video-camera with sampling rates of 200 f/s (HAS-220) and 2000 - 1000 f/s, respectively. These frames were captured and stored on a computer and used for further analyses (Movias Pro, NAC, Japan). Prior to all recording sessions, circle markers were placed on each rat and human’s skin at the vertebrae at L3 (a), hip (b), knee (c), ankle (d), and metatarsal joints (e), and the hip, knee, and ankle joints were defined as follows, hip joint: angle a-b-c, knee joint: angle b-c-d, ankle joint: angle c-d-e (Figure 1).
**2.3. Data Analysis**

Stick figures, which connected the circle marks on the skin and eye, ear and toe, were made from each frame. From these images, the onsets of swing (SW) and stance (ST) phases were determined by visual inspection. The hip, knee, and ankle joints were measured for the angle excursion graphs and angle-angle cyclographs. Raw kinematic data were averaged for ~10 step cycles. EMG data were synchronized with a high speed camera using a 200 Hz signal. Raw EMG bursts were rectified and averaged for 10 - 15 step cycles and averaged EMG waveforms were made for each muscle to compare the EMG activity patterns to the locomotor cycle. EMGs recorded from humans were also rectified and averaged. The angular excursion graphs and the duration of EMG bursts were demonstrated as % of cycle; onset of the SW phase (0%) and termination of the ST phase of the right hindlimb (100%). To detect commonly observed EMG activity patterns between Qp and Bp in a single subject, and between RBWM and humans for the same walking pattern, the qualitative EMG comparison was made. For this, we focused mainly on the temporal component such as the onset and/or the duration of EMG activity during a step.

**3. Results**

Both RBWM and humans could walk seamlessly on the treadmill belt. Figures 2 and 3 show serial drawings of a single step cycle of Qp and Bp walking in RBWM and humans, respectively. The tail of RBWM was not drawn in Figure 2.

**3.1. Angular Changes at the Hip, Knee and Ankle Joints in RBWM and Human**

Figure 4 shows representative serial angular excursions during three step cycles in the hip, knee and ankle joints of RBWM (A) and humans (B) during Qp (1) and Bp (2) walking on a level treadmill belt. All three joints in both RBWM and humans showed replicable across step cycles, but the angular excursion graphs of the same joint in the two groups were quite different.
Figure 3. Serial drawings of Qp (A) and Bp (B) walking in human subjects during a single step cycle. The numbers on each drawing indicate the order of the steps.

Figure 4. Representative three step cycle angular changes in hip, knee and ankle joints in RBWM (A) and humans (B) during Qp (1) and Bp (2) walking. Gait diagrams of the right hindlimb are shown at the bottom of each graph; the swing phase is shown by white bars and stance phase by gray bars.

3.1.1. Qp Walking in RBWM (Figure 4A-1)
Hip joint showed smooth flexion and extension transition during a single step cycle. The maximum flexion and extension were observed during the early and late ST phase, respectively. The mean mobile range of this joint was about 75° - 90°. From the initiation of the SW phase, knee joint continued to move toward extension until the transition period to ST phase and showed maximum extension. It then moved toward flexion until the end of the ST phase. The mean mobile range of this joint was about 50° - 100°. Ankle joint had a wider mobile range than other two joints throughout the single step cycle. From the late SW phase, this joint moved toward extension until the early ST phase. Then, it flexed for a short period and again moved toward extension until the late part of the ST phase, making two humps at the beginning (small) and the late part (large) of the ST phase, after which it started to flex toward the SW phase. The mean mobile range of this joint was about 50° - 105°.

3.1.2. Qp Walking in Human (Figure 4B-1)
From the end of SW phase, hip joint moved toward extension until the end of the ST phase, making a small hump at the beginning of this phase. It then rapidly flexed until the end of SW. The mean mobile range of this joint was 80° - 150°. Knee joint showed a basically similar angular excursion pattern to that of the hip joint. The maximum flexion and extension of this joint were observed at the mid to late SW phase and at the late ST phase, respectively, and these appeared earlier than those of the hip. At the end of the ST phase, it started quick flexion until the mid SW. The mean mobile range of this joint was about 80° - 145°. Ankle joint moved through a single step cycle similar to that during Bp walking. There were two humps, i.e., a small one at the early and a large one at the late phase of ST, and these were similarly observed in RBWM Qp walking. Between these humps during the ST phase, this joint first moved toward flexion until the mid ST phase and then extended until the end of the ST phase. The mean mobile range of this joint was about 85° - 110°.

3.1.3. Bp Walking in RBWM (Figure 4A-2)
Hip joint showed a small range of movement during a
single step cycle. At the end of the ST phase, this joint showed maximum extension and then continued to flex until the beginning of the next ST phase. The mean mobile range of this joint was about 85° - 105°, and this was as small as that during Bp walking. In contrast to the small movement range of the hip joint, knee joint showed saw-blades with clear extension and flexion during a step cycle. Throughout all phases of SW, this joint moved towards extension. The maximum extension of this joint was observed at the early phase of ST. Then, it showed gradual flexion until the end of the ST phase. At this timing, this joint showed maximum flexion at around 40°. The range of motion of this joint was from 40° to 80°.

3.1.4. Bp Walking in Human (Figure 4B-2)

From the late ST phase, hip joint moved towards flexion until the mid SW phase, followed by an extension until the end of the ST phase with a small hump at the beginning of the ST phase. This joint showed maximum extension and flexion at the end of the ST and at the end of SW phase, respectively. The mobile range of this joint was 150° - 180°. At the late phase of ST, knee joint started to flex and showed maximum flexion at the early phase of SW. It then rapidly moved toward extension until the end of the SW phase. The joint then remained at around 160° until the end of the ST phase. This constant joint angle period was not observed in RBWM, which showed a gradual decrease during this phase. The mobile range of this joint was 125° - 165°. Ankle joint showed similar angular excursion to that observed during Qp walking. From the end of the ST phase, this joint exhibited rapid extension until the mid SW phase. It then flexed until the end of SW and again started to extend during the early ST phase. Finally, towards the end of ST, it showed a gradual extension.

3.2. Angle-Angle Cyclographs of Knee-Hip and Ankle-Knee Relationships

The cyclographs show the average angle-angle relationships between two joints such as the knee and hip, and ankle and knee joints in RBWM (Figure 5) and humans (Figure 6) during Qp (A) and Bp (B) walking. In general, these figures showed that RBWM walked with their

![Figure 5. Representative averaged angle-angle cyclographs of hip-knee and knee-ankle joints in RBWM during Qp (A) and Bp (B) walking. For each group (A-1, A-2, B-1, B-2), the left and right side cyclograph shows the relationship between hip and knee, and knee and ankle, respectively. The upward and downward open triangles indicate the onset of the SW (gray line) and ST (black line) phase, respectively. The curved gray-colored arrows in the graphs show the direction of angle-angle joint movement.](image-url)
hindlimb joints rather flexed, while humans walked with their hindlimbs extended.

3.2.1. Qp Locomotion: Knee-Hip Joint Relationship (Figures 5A-1 vs 6A-1)

In RBWM, it was clearly observed that Qp walking was achieved with less hip joint movement. After the initiation of the SW phase, the hip joint started to flex, while the knee joint simultaneously started to extend, until the early ST phase. Then, the hip joint started to extend and the knee joint started to flex until the end of ST. The relationship between these two joints was observed linearly. In contrast to the hip joint of RBWM, that of humans showed a wider range of movement. After initiation of the SW phase, both hip and knee joints started to flex simultaneously. At the late SW phase, the maximum flexion of the hip joint and knee joint was observed. During this phase, the cyclograph curve was characterized by an “arc” shape. After that, both started to extend from the end of SW toward the late phase of ST. The maximum extensions of the hip and knee joints were observed during the ST phase. In the late ST phase, the knee showed rapid flexion, while the hip reversed from extension to flexion. The cyclograph moved in an anti-clockwise direction in both RBWM and humans.

3.2.2. Qp Locomotion: Ankle-Knee Joint Relationship (Figures 5A-2 vs 6A-2)

For RBWM, the cyclograph curve was characterized by a “crescent moon” shape during a single step cycle. At the late ST phase, the knee joint started to extend and the ankle started to flex, respectively. The maximum flexion of the ankle joint was observed at the late phase of SW. Then, both knee and ankle joints simultaneously started to extend until the early ST phase. After that, the knee joint continued to flex until the mid ST. The knee joint rapidly extended. At the late phase of ST, the knee and ankle started to flex and extend, respectively. For humans, the cyclograph was characterized by an “oval” shape. This is why both joints move in a similar way during a step cycle. After initiation of the swing phase, both started to flex and the curve moved in a clockwise direction. In this relationship, the cyclographs of both RBWM and human moved in a same clockwise direction.
3.2.3. Bp Locomotion: Knee-Hip Joint Relationship
(Figures 5B-1 vs 6B-1)
In RBWM, the cyclograph showed a “flat oval” shape with a longer horizontal (knee) axis. This shape was made by the rapid extension-flexion movement of the knee joint with less that of the hip joint. After initiation of the SW phase, the knee joint showed extension for this period, while the hip joint was weakly extended. Then, the knee joint started to flex until the late part of the ST phase, with weak hip extension. In humans, the cyclograph curve was characterized by a “triangle” shape during a one step cycle. This triangle shape cyclograph indicated that both the hip and knee joints showed similar flexion and extension cycles throughout a single step cycle. At the early SW phase, both hip and knee joints simultaneously flexed. During mid to late SW phase, the hip joint remained constant and the knee joint was extended. From the beginning of the ST phase, the hip joint showed gradual extension, while the extended knee joint was preserved. Then, at the late ST phase, both the hip and knee joints showed gradual flexion until the initiation of the SW phase. In both RBWM and human subjects, the cyclograph directions were anti-clockwise.

3.2.4. Bp Locomotion: Ankle-Knee Joint Relationship
(Figures 5B-2 vs 6B-2)
The cyclograph curve in RBWM was expressed by a “figure of eight”. During the SW phase, the knee joint rapidly extended, while the ankle joint showed gradual extension. After the onset of the ST phase, both the knee and ankle joint started to flex up to two-thirds of the ST phase. Then, the ankle began to extend, while the knee joint maintained flexion in the late ST phase. For humans, the cyclograph between the knee and ankle showed a “figure of eight” shape which was similar to that observed during RBWM Bp walking. At the early phase of SW, both the knee and ankle were flexed. Then, the knee joint was rapidly extended, while the ankle joint remained constant. Subsequently, both the knee and ankle started to flex at the beginning of ST. During the late ST phase, the knee showed flexion while the ankle showed extension.

3.3. EMG Burst Patterns between RBWM and Humans in Qp and Bp Walking

Figure 7 illustrates ensemble-averaged EMG waveforms through a single step cycle between different subjects in the same walking pattern. As in Figure 7, EMG components showing similar onset and/or duration between RBWM and humans during Qp and Bp walking are indicated by the black and white arrows in each averaged EMG.

3.3.1. Longissimus Lumborum
Although no common EMG bursts were observed in either RBWM or humans during Qp walking, a burst at the early ST phase was observed in both RBWM and humans during Bp walking.

3.3.2. Gluteus Medius
During Qp walking, this muscle showed small and large bursts at the end of SW and at the late phase of ST, respectively, in both RBWM and humans. During Bp
walking, there was a burst EMG activity at the mid-ST in both RBWM and humans.

3.3.3. Quadriceps
During Qp walking, two bursts were observed at the transition period from SW to ST and at the late phase of ST, however, the amplitudes were different.

3.3.4. Hamstrings
During Qp walking, the burst EMG activity was detected at the late ST phase in both RBWM and human subjects, while a burst component was found at the early phase of ST during Bp walking.

3.3.5. Gastrocnemius
In both RBWM and humans, there were burst activities at the late phase of ST. During Bp walking, an increased EMG activity was observed at the mid ST in both RBWM and humans.

3.3.6. Tibialis Anterior
This muscle showed a variety of activity patterns between RBWM and human subjects in either Qp or Bp walking.

3.4. EMG Activity between Qp and Bp Walking in RBWM and Humans

Figure 8 was prepared to detect common features of EMG activity during Qp and Bp walking in RBWM and humans, respectively. Overall, these results show that the muscle activities during the subject’s natural walking pattern were less active than those during the non-essential walking pattern, i.e. Qp < Bp for rat and Bp < Qp for human.

3.4.1. Longissimus Lumborum
In Qp-RBWM, there seem to be two burst activities during a single step cycle. They were observed at the transition period from the SW to ST phase and at the late part of ST. In Bp-RBWM, the EMG showed tonic activity during a step cycle with three burst activities. A burst EMG was obtained at the transition period from the SW to ST phases in both Qp and Bp walking. In human Qp, this muscle showed basically tonic activity with three bursts. One of these bursts was observed at the SW phase, and the other two were seen during the ST phase, respectively. The burst observed at the early ST phase during Bp walking was also obtained during Qp walking.

3.4.2. Gluteus Medius
In Qp-RBWM, this muscle showed small and large bursts at the end of SW and at the transition period from ST to SW phase, respectively. Except for these active periods, this muscle was rather silent. In Bp walking, this muscle showed three EMG bursts during a single step cycle. These were observed at the transition period from SW to ST, the mid-ST and the late ST phases. The burst at the late ST phase during Qp and Bp were similarly observed in RBWM. In humans, there were two burst activities and one burst activity during Qp and Bp walking, respectively. The onset timing and the duration were different.

3.4.3. Quadriceps
In RBWM, the EMG burst activities were different between Qp and Bp walking. During Qp walking, two burst activities were observed at the end of the SW (larger) and the mid-ST (smaller) phases. This muscle was activated at the end of ST during Bp walking. In human Qp, there were two bursts; one during SW and the other during the ST phase. A small burst was observed at the early phase of ST during Bp walking. This muscle showed different burst patterns and duration in humans.

3.4.4. Hamstrings
In RBWM during Qp walking, two EMG bursts were clearly observed at the late SW and at the end of ST phases, respectively. There were EMG bursts activated at the same timing during Bp walking. In humans, there were two EMG burst activities during Qp walking, while there was no clear burst during Bp walking.

3.4.5. Gastrocnemius
In Qp-RBWM, there were two bursts at the transition from SW to ST phases and during ST phase. On the other hand, there were three burst activities during Bp walking. These were observed at the transition period from the SW to ST, the mid-ST and the end of the ST phases. The first and the last ones were activated at the same timing in Qp-RBWM. In humans, two bursts were observed during Qp walking, while a single burst was seen during Bp walking. This burst was activated at the same timing during a step cycle (early phase of ST) during Qp walking.

3.4.6. Tibialis Anterior
In RBWM, this muscle showed a simple burst at the end of ST during both Qp and Bp walking. In humans, there were two burst activities in both Qp and Bp walking. The burst which appeared at the transition from the SW to ST phase was observed in both Qp and Bp walking.

3.5. Comparison of Averaged EMG Activity during Qp and Bp Walking in RBWM and Human

To compare each muscle activity during a single step...
cycle between Qp and Bp walking quantitatively, the area of averaged EMG activity of each muscle and the ratio between non-natural/natural walking, i.e. Bp/Qp in RBWM and Qp/Bp in humans was calculated (Table 1). The area of averaged EMG activity was defined as the shaded area shown in Figure 8 top-right. In both subjects, 5 of 6 muscles examined showed larger areas during non-natural walking than during natural walking, i.e. small EMG during Qp in RBWM and small EMG during Bp in humans. The highest ratio was observed in the muscles of H in RBWM and of Q in humans. The average values of area comparisons were 203.8 and 240.1 in RBWM and humans, respectively. This suggests that the EMG activity during non-natural walking was about two times higher than during each subject’s natural walking.

4. Discussion

This study examined the kinematic characteristics and spatiotemporal patterns of muscle activity during Qp and Bp walking in rats which had acquired the ability to walk bipedally and humans. We focused to show the locomotor and EMG similarities and differences observed between rats and humans during Qp and Bp walking. Our results show a number of similarities between rat and human walking, but they also revealed several features specific to humans.

4.1. General Differences during Qp and Bp Walking

We consider that the change in the skeletal system is one of the determining factors in the development of human like upright standing and Bp walking in humans. For both upright standing and the execution of Bp walking, it is necessary to bring the center of gravity (COG) close to the hip joint and acquire the ability to raise the upper part of the body (Figure 1B-2). To do this, humans have a unique skeletal system that is not observed in other four-legged animals. Such skeletal features including the lumbar lordosis, sacral kyphosis, round pelvis, and so on enable humans to walk bipedally against gravity [1]. For example, lordosis stabilizes the upper body over the lower limbs in humans by positioning the trunk’s center of mass above the hip. In human females, substantially proliferated lordosis is observed during pregnancy to

| Table 1. Comparison of averaged EMG activity (area) during Qp and Bp walking in RBWM (n = 5) and human subjects (n = 3). |
|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
|                                                   | RBWM                                           | Human subjects                                 |
|                                                   | Qp    | Bp    | Bp/Qp (%) | Qp    | Bp    | Qp/Bp (%) |
| L       | 22.8  | 58.2  | 255.3     | 101.7 | 66.9  | 152.0     |
| Gm      | 11.7  | 17.9  | 153.0     | 151.3 | 93.9  | 161.1     |
| Q       | 12.8  | 5.6   | 43.8      | 344.2 | 57.5  | 598.6     |
| H       | 8.3   | 44.0  | 530.1     | 155.3 | 67.2  | 231.1     |
| Gs      | 17.0  | 27.6  | 162.4     | 95.3  | 112.5 | 84.7      |
| TA      | 6.9   | 9.1   | 131.9     | 457.1 | 145.3 | 314.6     |
| Ave     | 13.3  | 27.1  | 203.8     | 217.5 | 90.6  | 240.1     |

Figure 8. Rearranged ensemble-averaged EMG waveforms through a single step cycle in RBWM (n = 5) and humans (n = 3) during Qp and Bp walking to compare the effect on the walking pattern in each subject. The arrows and the black bar have the same meaning as those in Figure 7.
compensate for the fetal load [7]. From these facts, it should be emphasized that these skeletal characteristics help such spatial body orientation for upright body posture. The typical Qp ungulates are not able to walk bipedally. This is because their COGs are always maintained at around the front of the animal’s body during walking and this position is clearly different from that of humans. Therefore, the bipedalism must strongly relate with the position of COG. In this sense, humans are a unique mammal which can adapt their Bp walking to various circumstances while maintaining the COG directly at the hip joint.

It is well known that primates can acquire the ability to walk bipedally and even clear obstacles on a walking path with motor learning [8-10]. This indicates that they are able to bring their COG near to the hip joint due to their acquired skeletal similarities to humans such as the lordosis in various situations [10-12]. However, their walking posture is not exactly the same as that of humans. This is due to the characteristic flexed body posture and called “bent-hip and bent-knee” walking posture [4]. As a result, their upper bodies are rather forward-bent. In our rat model, the rats could walk bipedally, but their body axis orientation was forwardbent, suggesting that the COG is not exactly on the hip joint (Figures 1A-2 and 2B). This walking posture surprisingly resembles to that of chimpanzees and other nonhuman primates. Why do they walk with such a posture? There are many kinds of four-legged mammals in nature and they have adapted their body structures suitable to their living environments during evolution. From this point of view, it is possible that their bodies are not suitable for erect posture due to skeletal constraints such as the different orientation of the ischium between primates and humans [13]. Substantially, the key to the skeletal structure for human-like walking posture can be found at around the pelvic girdle [14].

4.2. Motor Patterns during Qp Walking in Rats and Humans

Qp walking is a natural locomotor pattern for the rat, but it may be a challenging locomotor task for humans [15-17]. In this locomotor task, our findings revealed both similarities and differences between rats and humans. Among the similarities is the rhythmic and replicable hindlimb movement across consecutive step cycles which reflect the existence of a common rhythm generating system in both subjects [18]. The cyclographs showed that both knee-hip and ankle-knee relationships during Qp walking were in anti-clockwise and clockwise directions, respectively, in rats (Figure 5) and humans (Figure 6). These results suggest that the rhythmic hindlimb motion and its alignment were similarly controlled between rats and humans for the smooth and seamless execution of Qp walking.

We also found several differences in the mobile range of the hindlimb joint and the amplitude of each muscle during walking. The mobile ranges of the hip and knee joints during Qp walking in RBWM (Qp-RBWM) were smaller than that during Qp walking in humans (Qp-Human), and these two joints were generally flexed (Figure 4). On the other hand, the range of ankle in Qp-RBWM was larger than that in Qp-Human. Ankle plantar flexion during Qp-RBWM and Qp-Human was observed twice in a single step cycle. This angular excursion was well coupled with the EMG burst observed in Gs, which is the ankle plantar flexor muscle (Figure 4). The Q muscle in both RBWM and humans showed EMG bursts during the extension phase of the ipsilateral hindlimb knee joint and these bursts were observed during the SW phase of the same limb during Qp walking. Moreover, the burst activities observed in Gm and H were common between RBWM and humans, suggesting the proximal muscles are also used in a similar manner in both rats and humans (Figure 7). These results suggest that both subjects recruited common motor patterns in the hindlimbs during Qp walking, while having their own mobile range of hindlimb joints.

4.3. Motor Patterns during Bp Walking in Rats and Humans

RBWM showed stable BP walking, but the Bp walking pattern was not exactly the same as that of humans (Figure 2B). In this locomotor task, a clear difference was observed at the shortened period of the SW phase in RBWM. The ratio between the period of the SW/ST phases during Bp walking was about 1:9 in RBWM and 3:7 in humans; however, the ratio itself was preserved with replicable angular excursions which reflect the existence of a rhythm generator in rats and humans (Figure 4, [18]).

Conversely to Qp walking, many muscles were highly activated during Bp walking in RBWM (Figures 7 and 8). However, it should also be indicated that all of these muscles were phase-dependently activated [16]. For example, in Q muscle during Bp walking, the activity was transiently observed from the end of ST to the SW phases and this was for both extension of knee joints and flexion of hip joints in RBWM. In contrast to the RBWM Q EMG activity, that of humans was seen from the end of SW to the mid-ST phase, and seems to be used for knee extension and maintenance. In fact, Q is a bi-functional muscle used to flex the hip and extend knee joints, and these results suggest that both rat and human subjects could use this muscle for Bp walking. The activity of the H muscle in Bp-RBWM was maintained throughout the
ST phase (early > late), while human H muscle showed a different activity pattern during the same period. This H muscle activity and the lower activity of Q muscle caused knee flexion in RBWM during the ST phase. This knee flexion lowered the COG of Bp walking rats and may have forcibly terminated stable Bp walking on the treadmill belt. Certainly, the knee joint gradually flexed during the ST phase in RBWM (Figure 4A-2). To correct this walking posture, it is necessary to bring the lowered COG higher to the proper level. The small EMG burst activity of the H muscle observed at the end of the ST phase in the ipsilateral limb must be recruited for this. Continuous knee flexion during the ST phase also forced initiation of the SW phase of the contralateral limb and the short SW phase was indispensable for the next step cycle. It was interesting that the activity pattern of this H muscle was not exactly the same between RBWM and human subjects during Bp walking, but the largest EMG amplitude of this muscle was observed at the transition from the SW to ST phase. The same EMG activity component is present in other muscles such as the L, Gm and Gs. Four of six muscles examined, showed surprisingly similar activation patterns and/or components during Bp walking in both rats and human subjects. Moreover, three of these muscles are activated in both Qp and Bp walking. These results may indicate that both subjects have a similar control system for the different walking patterns.

4.4. Motor Patterns between Qp and Bp Walking in Rats and Humans

Between Qp and Bp walking, we also found many similarities and differences in the characteristic EMG bursts in both RBWM and human subjects, respectively (Figure 8). The Gm muscle is the extensor of the hip joint. The largest EMG burst of this muscle was mainly observed at the end of the ST phase in both locomotor patterns. This reflected that the ipsilateral hindlimb is pushing the walking surface before toe off. The hip joint excursion in both Qp and Bp walking was quite small (Figure 4) and this seems to reflect the smaller EMG burst activity of this muscle in RBWM. During Bp walking, the other two bursts were observed at the beginning of ST and the mid-ST phases and were probably recruiting for the extension of the hip joint and the maintenance of its joint angle during Bp walking. In humans, the EMG activities of the Gm muscle in both Bp and Qp walking resemble those of the Q muscle. This suggests that these two muscles are co-functioning in humans. In Bp walking, Gm muscle bursts observed during the ST phase extended the angle of the hip joint from 165° to 180°. It was noteworthy that the Gm EMG burst was terminating when the ipsilateral knee started to flex together with hip flexion. On the other hand, during Qp walking, Gm EMG bursts were observed twice at the end of SW and the ST phase, respectively. This pattern was similarly observed in the Q muscle, suggesting the Gm muscle co-activates with the Q muscle. The latter activity was observed during extension of the ipsilateral hip joint. The former was observed when the hip joint showed flexion and the knee joint started to extend. It seemed unlikely to be related to hip joint extension, but this might be a characteristic EMG burst observed when human subjects are walking on a treadmill belt quadrupedally (hand-foot crawling).

A clear difference was also observed in the Q muscle in humans. The burst of this muscle mainly corresponded to the flexion of the knee joint. It is considered that the Q muscle EMG burst during the ST phase of the ipsilateral hindlimb inhibits excessive knee flexion during Qp walking. EMG bursts of Q in Bp-human were small and mainly observed during early ST phase. During Qp walking, the Q EMG burst was also observed at the ST phase and had large amplitude. However, the function of this EMG burst during Qp walking seems to be different. The EMG burst during Bp maintained the angle of the knee joint at around 165°, while Q EMG bursts during the ST phase in Qp walking were observed during the extension phase of the ipsilateral knee joint. These facts indicate that the human subjects could appropriately recruit each muscle’s functional role for different walking patterns. Between Qp and Bp walking, five of six and three of six muscles showed similar EMG burst activities in rats and humans, respectively. In rats, this result suggests that their Qp and Bp walking is achieved by similar muscular activations. In contrast to the results from rats, humans tend to use each muscle selectively to fit different walking patterns. This may be due to the recruitment of voluntary control in Qp walking that is not a natural pattern for adult humans [16].

4.5. Expected Energy Consumption between Qp and Bp in Rats and Humans

For rats and adult humans, the Bp and Qp walking patterns were not natural, respectively. The general high activation of each muscle during Bp-RBWM and during Qp-Human in the present study implies that RBWM and humans need a lot of energy during their Bp and Qp walking, respectively. In fact, the EMG amplitude during Qp-RBWM was smaller than that during Bp-RBWM, and that during Bp-human was smaller than that during Qp-human (Figure 8). It is not possible to calculate energy consumption from EMG activity, however, it might be possible to argue that the increased activation should increase the cost. From this point, we had calculated the area of averaged each muscle EMG activity as shown in Figure 8 (top right dark shaded area).
results are summarized in Table 1. Five of six muscles in both RBWM and humans showed about a 1.5 to 6 times larger area in Bp and Qp walking, respectively. The average comparison showed about a 2 to 2.4 times larger area in the non-natural walking patterns. Although the areas of Q in RBWM andGs in humans were smaller during the non-natural walking patterns, the majority of the muscles that underwent EMG recording showed similar trends. This suggests that the non-natural walking patterns was made by a lot of efforts and probably required more energy than for the natural walking patterns. The smaller areas in Q and Gs, however, indicate that these are probably not necessary or are less affected in both Bp and Qp in RBWM and humans. This is because a clear EMG burst was observed in both muscles and this burst induced clear joint movement during different walking patterns. Nakatsukasa et al. [19,20] showed that Bp-trained nonhuman primates consume less metabolic energy during their natural Qp walking than Bp. This suggests that RBWM Bp and human Qp walking may demand more precise neuronal control, such as cortical control, for unfamiliar body posture requirements and/or intersegmental coordination as compared to their habitual quadrupedal and bipedal walking patterns.

As described above, energy consumption is the one of the key determinants in the selection of a walking pattern for animals [21]. Sockol et al. [13] investigated the cost of transport in five chimpanzees between Qp and Bp walking. They found that an individual comparison showed a significant difference between Qp and Bp walking in both juvenile males (n = 2) and adult females (n = 3). For two juveniles, the cost of transport was significantly lower during Qp walking and one of the adult chimpanzees showed a similar result. Surprisingly, one of the other two female adults showed a significantly lower cost of transport during Bp walking (Bp lower energy consumption chimpanzee). They also compared the kinematic characteristics and found that the Bp lower energy consumption chimpanzee extended her hips and knees to a similar degree during Qp and Bp walking, while the others showed more flexed limb orientation during Bp walking. Even this chimpanzee, however, bent her body axis rather forward and this suggests that the body axis, or at least the hip and knee joints, functionally interlock during walking as in a hinge. The anatomical difference in ischium between chimpanzee and human is clear; the former is oriented distally, while the latter is dorsally oriented. This spatial difference reduces the ability of the hamstring, which produces an extensor moment when the femur is extended, relative to the pelvis in chimpanzees. This biomechanical constraint seems to be one of the main reasons preventing chimpanzees from walking like humans. From these, it was also suggested that slight kinematic changes induce large differences in walking cost and are consistent with previous studies showing that differences in posture can affect cost [22, 23]. For pregnant human females, the fetal load moves her COG forward and thus bends her posture forward. This destabilized Bp standing posture is usually compensated by enhanced lumbar lordosis and translating the COG backward [7]. Without this skeletal compensation, such an instable posture must be counteracted by the back and proximal lower limb muscles, requiring much more energy than a stable posture. This is reasonable because the energy consumption must be smaller when the COG is on the hip joint than when it is far from the hip. From these facts, it is clear that the skeletal system also affects the energy consumption during walking for individual animals.

5. Conclusion

Overall, this study provides new data on EMG activities and kinematics during Qp and Bp walking in both rats and humans. Rats that acquired Bp walking capability and humans showed several common kinematics and EMG burst patterns during the same walking task. This suggests that they share common neuronal control systems, at least at the spinal cord level. The differences in posture during Bp walking may be due to the skeletal constraints that do not allow four-legged mammals including nonhuman primates to extend their hip and knee joints and even the body axis. Due to this skeletal characteristic, the elaboration of Bp walking requires a lot more energy than for Qp walking in rat.

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