

The Lower Limb and Mechanics of Walking in *Australopithecus sediba*

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The discovery of a relatively complete *Australopithecus sediba* adult female skeleton permits a detailed locomotor analysis in which joint systems can be integrated to form a comprehensive picture of gait kinematics in this late australopith. Here we describe the lower limb anatomy of *Au. sediba* and hypothesize that this species walked with a fully extended leg and with an inverted foot during the swing phase of bipedal walking. Initial contact of the lateral foot with the ground resulted in a large pronatory torque around the joints of the foot that caused extreme medial weight transfer (hyperpronation) into the toe-off phase of the gait cycle (late pronation). These bipedal mechanics are different from those often reconstructed for other australopiths and suggest that there may have been several forms of bipedalism during the Plio-Pleistocene.

The locality of Malapa, South Africa, has yielded two relatively complete skeletons of *Australopithecus sediba*, dated at 1.977 million years ago (1, 2). This species has a combination of primitive and derived features in the hand (3), upper limb (4), thorax (5), spine (6), and foot (7) in a hominin with a relatively small brain (8), a human-like pelvis (9), and a mosaic of *Homo*- and *Australopithecus*-like craniodental anatomy (1, 10, 11). The foot in particular possesses an anatomical mosaic not present in either *Au. afarensis* or *Au. africanus* (7), supporting the contention that there were multiple forms of bipedal locomotion in the Plio-Pleistocene (12). The recent discovery of an *Ardipithecus*-like foot from 3.4-million-year-old deposits at Burtele, Ethiopia, further shows that at least two different kinematic solutions to bipedalism coexisted in the Pliocene (13). Here we describe the lower limb of *Au. sediba* [specimen numbers and attributions are provided in table S1 (14)] and propose a hypothesis for how this late australopith walked.

Lower Limb of MH1

The holotype of *Au. sediba* is Malapa Hominin 1 (MH1), a juvenile male partial skeleton whose lower limb consists of a right proximal femur (fig. S1), small shaft fragments from the tibia and fibula, and foot bones already described (7) (table S1). The proximal femur is australopith-like, with a long, anteroposteriorly compressed

femoral neck (fig. S2) and low neck-shaft angle (110° to 115°) (table S2). Posterolaterally, there is a third trochanter, inferior to which is a well-developed hypotrochanteric fossa, a human feature reflecting a large insertion area for the gluteus maximus (15).

Lower Limb of MH2

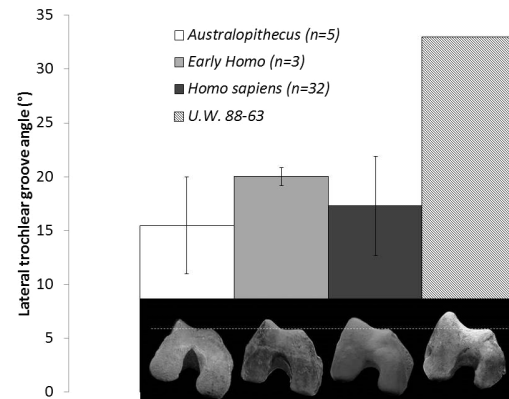
The lower limb of MH2, an adult female, consists of a right femoral head and neck, part of the proximal femoral shaft, the left proximal fibula, and the right knee joint (including the patella). Additionally, as already described (7), MH2 preserves an articulated distal tibia, talus, and calcaneus and a partial fifth metatarsal. The proximal femur preserves much of the head and neck (fig. S1). As in MH1, the neck is anteroposteriorly compressed (table S2). Viewed superiorly, the head appears to be prolonged anteriorly, as is the case in most humans (16).

The right knee of MH2 is represented by an 86.7-mm-long fragment of the distal femur (fig. S3), two fragments that conjoin to form most of the tibial plateau (fig. S4), and a relatively complete patella composed of two conjoining frag-

ments, one of which remains partially embedded in calcified sediment. This part of the patella has been digitally extracted from micro-computed tomography (μ CT) scans, and a nearly complete knee cap has been reconstructed (fig. S5). The posteromedial part of the distal femur has been sheared away, but the lateral condyle, patellar articular surface, distal shaft, and most of the medial condyle are well preserved. The bicondylar angle is estimated to be ~9°, which is within the range of modern humans but is low for an australopith (table S3). There is a sutured hollow just superior to the patellar surface, evidence of contact with the patella in a fully extended position (17) (fig. S6). There is a strong medial condylar boss, an anatomy unique to hominins, and evidence for a “tibial dominant” knee capable of full extension (18) (fig. S7). Most notable is the high lateral patellar lip. The lateral trochlear groove angle (19) is 31.3°, 3 SD above the modern human mean (20) and far greater than any ape trochlear angle, because apes tend to have flat trochlear grooves (Fig. 1). A high lateral patellar lip serves as a bony mechanism for patellar retention during bipedal gait (21, 22). The extension of the lateral lip in *Au. sediba* is not a function of overall anterior expansion of the patellar surface, as is found in *Homo* (18), and is restricted just to the lateral side (fig. S8). Laterally, there is a deep groove for the popliteus, an internal rotator of the tibia and stabilizer of the knee. This muscle may have been important in resisting internal rotation of the femur during stance phase. The relatively narrow tibial spines on the MH2 tibial plateau suggest enhanced knee mobility (23), although this anatomy may also be related to the small size of *Au. sediba* (24).

On the proximal tibia, the medial condyle is flat and the lateral condyle is slightly convex anteroposteriorly, similar to the condition found in other small australopiths (specimens A.L. 129-1 and StW 514), although the functional importance of this convexity is unclear (25). There appears to be a small notch on the lateral plateau, perhaps indicating the presence of a double meniscus attachment and thus possibly greater osteoligamentous

Fig. 1. The lateral patellar lip. The lateral trochlear groove angle (19) is similar in *Australopithecus* (TM 1513, Sts 34, A.L. 129-1, A.L. 333-4, and A.L. 333w-56), early *Homo* (KNM-ER 1472, KNM-ER 1481, and KNM-WT 15000), and modern humans (20). This measurement in MH2 (U.W. 88-63) is over 3 SD higher than in modern humans. Apes have flat trochlear grooves (18) (fig. S7) and thus lateral trochlear groove angles near zero. Bottom images, from left to right, are as follows: TM 1513, KNM-ER 1472, modern human, and MH2, all scaled to the same size. They have been positioned so that the medial patellar surface is horizontal, which corresponds closely to the orientation recommended in (18). Note the extreme lateral patellar lip in MH2.



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control over rotation at the knee as in *Homo* proximal tibiae (26). However, the absence of this notch in other australopiths does not necessarily imply the absence of a double insertion (27).

The patella is small, measuring 27.1 mm wide mediolaterally and 13.1 mm thick anteroposteriorly. It is 24.7 mm tall superoinferiorly, which is probably just short of the actual height, because there is some damage to the distolateral aspect of the apex. The posterior part of the patella is human-like in being strongly convex mediolaterally (fig. S9), with a high central keel separating the condylar facets medially and laterally.

The most proximal 97.1 mm of the left fibula of MH2 is preserved as four conjoining fragments (fig. S10). The fibula is more gracile than modern

ape fibulae and in this respect resembles specimen OH 35 (fig. S11). The MH2 fibula has an osteophytic growth at the biceps femoris insertion.

Lower Limb of MH4

MH4, an adult or near-adult individual of unknown sex, is represented by two conjoining pieces of a tibia described previously (7). Here we provide an estimated total length, possible because the proximal tibia, though not recovered, has left a natural cast of its anterior surface in the calcified sediment (fig. S12). We estimate total tibial length at approximately 271 mm, with a possible range of 267 to 275 mm, depending on the degree of proximal tibial retroversion and the proximal projection of the tibial spines.

Fig. 2. Hyperpronation.

(A) The pedal bones of *Au. sediba* are superimposed on a human foot in dorsal view. These bones are not all from the same individual (see table S1 for details). We hypothesize that MH2 would have contacted the ground along the lateral edge of an inverted foot. This would generate a ground reaction resultant (blue arrows) that would be positioned lateral to the joints of the foot, creating a large pronatory torque (red arrows). Although apes often land along the lateral edge of an inverted foot (30, 31), they swing their body mass laterally over the stance foot during bipedal gait, effectively producing a counteracting supinatory torque. *Au. sediba* had a pelvis with sagittally oriented ilia (9), suggesting a human-like abductor mechanism and in turn suggesting a medially positioned center of mass relative to the stance leg (illustrated by the large blue arrow at bottom left). This position of the center of mass would exacerbate the pronatory torque at the subtalar, midtarsal, and tarsometatarsal joints. (B) Excessive pronation on a weight-bearing foot (curved red arrow) causes a chain of rotatory movements proximal to the foot. The tibia internally rotates (green arrow) as the talus plantarflexes and adducts. The femur also internally rotates [(42, 43); curved blue arrow], increasing the lateral deviation of the patella (small blue arrow). Pronation at the foot causes an anterior pitch of the center of mass [(34, 43); black arrow], counteracted by hypertordosis of the lumbar region (6). *Au. sediba* possesses anatomies that are adaptive to, or consequences of, these motions.

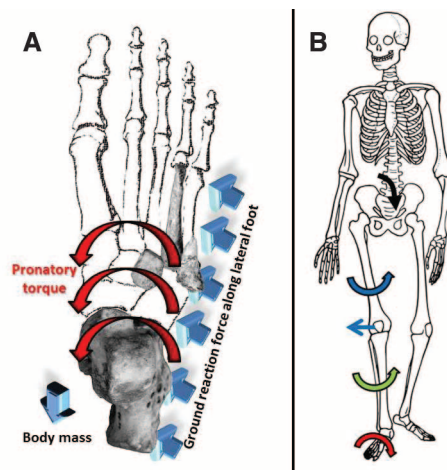
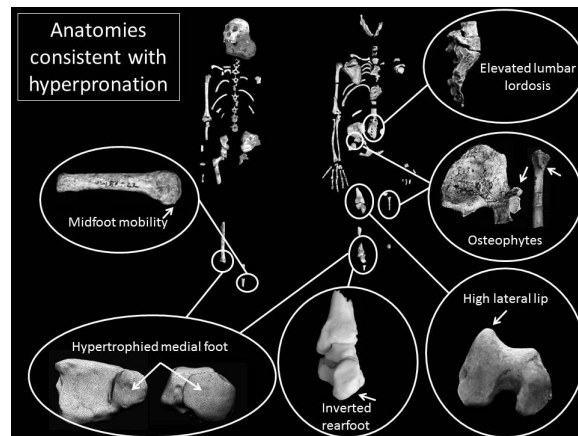


Fig. 3. Skeletons of *Au. sediba*:

left, MH1 (pictured with MH4 tibia); right, MH2. Several anatomies of these skeletons are consistent with a hyperpronating gait. The base of the fourth metatarsal is convex, indicating that the midfoot was hypermobile. The medial malleolus and talar head are hypertrophied, reflecting loading of an inverted foot and mobility at the talonavicular joint, respectively. The rearfoot is in an inverted position, a risk factor for hyperpronation in modern humans. Hyperpronators internally rotate the femur and tibia and are at a greater risk for patellar dislocation.

The high lateral patellar lip reduces the risk of patellar subluxation. Osteophytes on the origin of the rectus femoris/iliofemoral ligament proximal attachment and insertion of the biceps femoris indicate soft tissue strain: possible consequences of a hyperpronating gait. MH2 has elevated lumbar lordosis (6), perhaps to compensate for the excessive anterior pitch of the center of mass common in hyperpronators.



The Kinematics of Walking in *Au. sediba*

The anatomy of the foot (7), spine (6), pelvis (9), and knee (this paper) indicate that *Au. sediba* was an obligate biped. Based primarily on the lower limb, pelvic, and vertebral morphology of MH2, and to a lesser extent on the pedal morphology of MH1, we propose that *Au. sediba* was a hyperpronator (28) with exaggerated medial weight transfer during the stance phase of terrestrial bipedalism (Fig. 2). Modern human hyperpronators serve as a kinematic reference model for this gait and its musculoskeletal consequences. We suggest that MH2 expressed this kinematic variation, and *Au. sediba* did with regular frequency, although the hypothesis that the entire species walked in this manner will require further testing with additional fossil material (14).

At heel strike of bipedal locomotion, humans commonly have a slightly supinated rearfoot and forefoot, which are passively driven by the ground reaction force into pronation of the subtalar and more distal joints during the subsequent mid-stance phase of walking. Video and plantar pressure data reveal that apes contact the ground with the heel (29) and often the lateral midfoot simultaneously in what has been termed “inverted heel-strike plantigrady” (30, 31). The abducted hallux serves a stabilizing role during quadrupedal walking in apes, contributing little to propulsion (31). The divergent hallux is suggested to have served a similar stabilizing role during bipedal walking in *Ardipithecus ramidus* (32).

In a small percentage of modern humans, the foot is excessively inverted (termed forefoot and/or rearfoot varus) during the swing phase of walking, resulting in heel strike with the foot in a highly supinated posture, with ground contact established along the lateral border of the heel and forefoot (33). Contact between the ground and the lateral side of the foot introduces a large pronatory torque around the subtalar and more distal joints, which drives the foot into pronation (33, 34). As the foot is driven into pronation, there are high medially directed torques that can not only cause excess loading on the bones of the medial column of the foot (35) but also stress the soft tissues, such as the ligaments supporting the medial longitudinal arch and the muscles whose tendons insert plantomedially, particularly the tibialis anterior and tibialis posterior. Plantar fasciitis, medial tibial stress syndrome (shin splints), and tibial stress fractures are therefore common injuries experienced by late and hyperpronating modern humans (34). Although hyperpronation can have pathological consequences in modern humans, we are proposing here that the skeleton of *Au. sediba* reveals a suite of anatomies that are adaptive for, or consequences of, this kind of walking (Fig. 3).

Reconstruction of the conjoined elements of the rearfoot of MH2 demonstrates that the calcaneus had an inverted set (fig. S13), which is a contributing factor to hyperpronation in modern humans (36). However, the inverted heel itself

would not necessarily produce a pronatory torque, because there is considerable variation in the position of the subtalar joint axis relative to the ground reaction force location (37). MH2 also had a gracile calcaneal tuber (7), with a superiorly positioned lateral plantar process, which reduced the surface area of the plantar aspect of the heel in *Au. sediba* and would have reduced the effectiveness of the subcalcaneal heel pad (based on size information rather than material properties), which has been shown to dissipate peak stress during heel strike (38). To compensate, we hypothesize that *Au. sediba* landed simultaneously on the heel and along the lateral foot at touch down, in much the same way that African apes walk (inverted heel-strike plantigrady) (31). This is achievable in a bipedal hominin that has full knee extension by slightly increasing normal plantarflexion angle during foot contact. Because of the wider midfoot and forefoot, landing along the lateral side of an inverted foot would provide a large moment arm around the midtarsal and tarsometatarsal joints that would also transfer to the subtalar joint. Thus, a large

pronatory torque would drive the foot into pronation (Fig. 2). There is suggestive evidence for excessive pronation in the *Au. sediba* tarsals. An elevated degree of pronation is possible in *Au. sediba*, because the subtalar joint has a high radius of curvature and is therefore quite mobile and capable of an extreme range of motion (7). The relatively large talar head of MH2 (7) may signal elevated talonavicular mobility, especially because this joint is central to midfoot pronation in humans (39).

Landing on an inverted foot would also load the medial portion of the tibiotalar joint and introduce a shear force across the medial malleolus. This may explain the form of the medial malleoli of both MH2 and MH4, which are mediolaterally thicker than those of other fossil hominins or modern humans (7). However, pronation does not occur at the tibiotalar joint but at the subtalar joint and joints of the midfoot. As the foot is driven into pronation by a high pronatory torque, the more distal parts of the medial foot would be excessively loaded (35). A foot adapted for this kind of locomotion may therefore be expected to

exhibit increased mechanical reinforcement of bones in the medial portion of the foot. We predict that, if additional foot elements are recovered, we will see greater joint and diaphyseal robusticity in medial relative to lateral tarsals, metatarsals, and phalanges (Table 1).

During midstance, the foot is more mobile and better able to conform to its substrate. In hyperpronators, the talus adducts and plantarflexes excessively, dropping the longitudinal arch and contributing to hypermobility of the midfoot. Although we hypothesize that *Au. sediba* possessed an arched foot (7) (fig. S14), there is also evidence for midfoot mobility. A right fourth metatarsal, possibly from MH1, has a highly convex base dorsoplantarily, suggesting the presence of midfoot flexion or a “midtarsal break” (40). Other hominin fourth metatarsal bases from *Au. afarensis*, *Au. africanus*, and the OH 8 foot are human-like and do not exhibit evidence for a midtarsal break (Fig. 4) (40, 41). The convexity of the Malapa fourth metatarsal is thus unexpected and implies more mobility at the lateral tarsometatarsal joint in this hominin than in any other. We suggest that the seemingly contradictory anatomies in the foot of *Au. sediba* (possession of an arched foot and long plantar ligament together with midfoot mobility) can only be explained in the context of a bipedal foot that hyperpronates when weight-bearing (14).

In modern humans, excessive pronation may have damaging effects in lower limb joints proximal to the foot. In hyperpronators, the tibia and femur both internally rotate excessively (42, 43) under a patella that is relatively fixed by the rectus femoris attachment to the anterior inferior iliac spine (AIIS) as the quadriceps femoris contracts to extend the leg during toe-off (34). Because this occurs late in the gait cycle, during knee extension, the patella is pulled laterally, and thus hyperpronators are at risk for both patellofemoral pain (44) and patellar subluxation (45, 46). A hominin bony adaptation that helps prevent patellar subluxation is a raised lateral lip of the distal femur (21, 22). The extreme lateral patellar lipping of MH2 (Fig. 1) (figs. S7 and S8) may be an adaptation to resist injurious lateral translation of the patella during hyperpronation of the foot and resulting internal rotation of the tibia and femur during late stance phase. This skeletal adaptation in *Au. sediba* may also implicate a reduced or absent vastus medialis obliquus in counteracting lateral translation of the patella. The fact that a lateral lip is present at birth (17, 18) can be extrapolated to indicate that the species *Au. sediba* (and not just MH2) was adapted for this kind of locomotion (14). Furthermore, the large popliteal groove present on the MH2 femur may indicate strong muscular involvement in counteracting the internal rotation of the femur on a fixed tibia, because the popliteus acts as an external rotator of the femur during stance phase.

Hyperpronation drives the entire leg medially during stance phase and may strain any muscle

Table 1. Evidence for hyperpronation in *Au. sediba*.

Hyperpronating biomechanics	Anatomical predictions	Morphology in <i>Au. sediba</i>
Initial ground contact on an inverted foot, resulting in high medially directed forces on tibiotalar joint	Inverted calcaneus and thick medial malleolus	Calcaneus in inverted set and predicted forefoot in varus set; thickest medial malleoli of any known hominin.
Excessive pronation at subtalar joint	Increased mobility at subtalar joint	High radius of curvature of talar facet on calcaneus.
Excessive pronation at midtarsal joints and tarsometatarsal joints	Mobile midfoot in coronal plane	Greatly enlarged talar head, suggestive of talonavicular mobility.
Increased strain on soft tissue of medial foot	Components of medial arch, plantar aponeurosis (if present), and tibialis posterior tendon under stress	Currently unknown. Predicted robust navicular tuberosity; reduced metatarsophalangeal joint extension if plantar aponeurosis taut.
Increased strain on foot bones distally and medially	Increased robusticity of medial tarsals, metatarsals, and phalanges	Currently unknown. Predicted to be relatively robust medially; if partially divergent, hallux could also help counter pronatory torque.
Lowered arch and increased midfoot mobility	Increased sagittal plane dorsiflexion evident in bones of midfoot	Convex surface to base of fourth metatarsal indicative of midtarsal break; predicted concavity of metatarsal facets on cuboid.
Increased knee mobility	Greater rotatory capacity and greater role for knee stabilizers, such as popliteus and biceps femoris	Tibial spines close together and enlarged popliteal groove on distal femur.
Increased internal rotation of femur	Increased strain on muscles crossing both hip and knee joints	Osteophytic growths on both origin for rectus femoris and insertion of biceps femoris. Predicted to have enlarged origin on the anterior superior iliac spine for sartorius.
Increased risk of patellar subluxation	Bony adaptations for patellar retention	Highest lateral patellar lip of any known hominin.
Anterior tilt of the pelvis	Increased lumbar lordosis	Last lumbar vertebra has very high wedging angle.

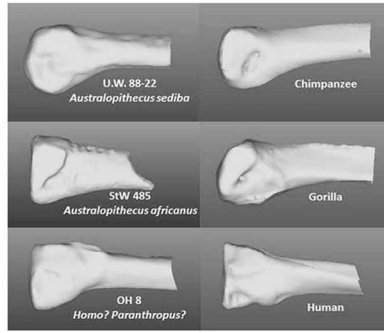
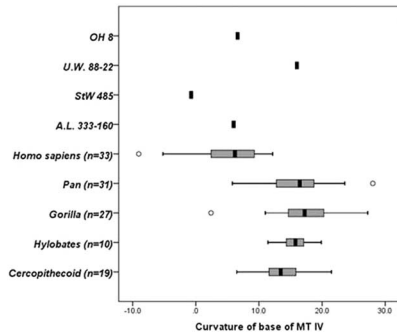


Fig. 4. The fourth metatarsal. (Left) Curvature of the base of the fourth metatarsal in fossil hominins, humans, apes, and monkeys (40). Humans and most fossil hominins, including *Au. afarensis* (41), have flat bases, consistent with a stiff and immobile midfoot. The *Au. sediba* fourth metatarsal has a convex base. [The box-and-whiskers plot shows the median (vertical line), upper and lower quartiles (box), range (whiskers), and outliers (circles) for each group.] (Right) Digital renderings of the fourth metatarsal in medial view, illustrating the dorsoplantarly curved base morphology in apes and *Au. sediba* and the flat base in other hominins and in modern humans.

crossing both the knee and the hip joints (e.g., the rectus femoris), particularly those inserting laterally in the leg (e.g., the long head of the biceps femoris). As previously mentioned, the insertion for the biceps femoris on the proximal fibula is osteophytic, indicative of elevated strain on this insertion area (fig. S10). The MH2 ilium has an unusually large and projecting AIIS (fig. S15), suggesting that the rectus femoris (and/or iliofemoral ligament) was under considerable strain during gait. As the femur internally rotates and adducts, the AIIS will be stressed by excessive stretching of the rectus femoris tendinous origin. Finally, hyperpronators experience an anterior pitch of the center of mass relative to the hip joint (43), requiring compensatory hyperlordosis to shift the center of mass posteriorly back over the hip joints, often resulting in lower back pain (34). The last lumbar vertebra of *Au. sediba* has very high dorsal wedging, suggesting elevated lordosis (6), which may have adapted this species to the challenges of being a hyperpronating biped.

Although we find the evidence compelling that *Au. sediba*, or at least MH2, was a hyperpronating biped (Fig. 3 and Table 1), the selective advantage of this form of bipedality is unclear. There is little evidence that other known australopiths were hyperpronators, because the peculiar anatomies of the *Au. sediba* foot, knee, and hip are not found in earlier australopiths. Recent work on the Laetoli footprints (47, 48) suggests that although the makers of the prints (presumably *Au. afarensis*) walked with a human-like gait, they had slightly less medial weight transfer. The hallux of *Au. afarensis* is domed and robust (49), indicating that weight transfer was more human-like than ape-like, but there probably were at least subtle differences in how *Au. afarensis* walked as compared to most modern humans. We hypothesize that terrestrial bipedalism in *Au. sediba* also differed subtly from that in most humans today, with *Au. sediba* engaging in more weight transfer on to the medial foot (hyper-

pronation) rather than less, as may have been the case with *Au. afarensis*.

Our interpretation of Malapa skeletal morphology extends the variation in *Australopithecus* locomotion. As suggested by others (7, 12, 13), there were different kinematic solutions for being a bipedal hominin in the Plio-Pleistocene. The MH2 skeleton provides insight into one of those potential solutions: hyperpronation. This mode of locomotion may be a compromise between an animal that is adapted for extended knee bipedalism and one that either still had an arboreal component or had re-evolved a more arboreal lifestyle from a more terrestrial ancestor. There is some postcranial evidence that the South African species *Au. africanus* may have been more arboreal than the east African *Au. afarensis* (50, 51), and a hypothesized close relationship between *Au. africanus* and *Au. sediba* (1, 11), along with features in the upper limbs of the latter thought to reflect adaptations to climbing and suspension (3, 4), is consistent with a retained arboreal component in the locomotor repertoire of *Au. sediba*. Pronation is an important foot motion that shifts weight onto the medial side of the foot in climbing apes (52, 53) and serves a role in weight transfer, shock absorption, and negotiation of uneven substrates during human bipedal gaits. An animal that was adapted to do both reasonably well may have had to support an increasingly mobile foot by evolving a large mobile medial column and important stabilizing anatomies at the knee and hip, in order to survive in these dual worlds.

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28. Pronation of the foot is a triplanar motion (eversion, abduction, and dorsiflexion occurring simultaneously) and is a normal function of bipedal foot mechanics in order to absorb ground reaction forces and accommodate uneven substrates. It converts the foot from a rigid structure at heel strike to a more mobile one at midstance. Hyperpronation is poorly defined in a clinical sense, because there is no distinct cutoff between those that excessively pronate and those that do not. However, here we define it as continued pronation into the part of stance phase when the foot should be resupinating (usually at the later part of midstance and during the entire propulsive phase). Extended (timing-wise) pronation is caused both by the magnitude of pronation (in degrees) that occurs as a result of the large pronatory torque generated by contacting the ground on a varus heel and forefoot, and the timing of foot motion. Because of the excessive motion occurring at the subtalar, midtarsal, and tarsometatarsal joints, the foot fails to resupinate at late stance and pushoff.
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Acknowledgments: We thank the South African Heritage Resource agency for the permits to work at the Malapa site; the Nash family for granting access to the Malapa site and continued support of research on their reserve; the South African Department of Science and Technology and the African Origins Platform (AOP), the South African National Research Foundation, the Evolutionary Studies Institute, the University

of the Witwatersrand, the University of the Witwatersrand's Vice Chancellor's Discretionary Fund, the National Geographic Society, the Leakey Foundation, the Palaeontological Scientific Trust, the Andrew W. Mellon Foundation, the Ford Foundation, the U.S. Diplomatic Mission to South Africa, the French Embassy of South Africa, the A.H. Schultz Foundation, Boston University, Duke University, a Ray A. Rothrock '77 Fellowship and International Research Travel Assistance Grant of Texas A&M University, and the Oppenheimer and Ackerman families and Sir Richard Branson for funding; the University of the Witwatersrand's Schools of Geosciences and Anatomical Sciences and the Bernard Price Institute for Palaeontology for support and facilities; the Gauteng Government, Gauteng Department of Agriculture, Conservation and Environment and the Cradle of Humankind Management Authority; and our respective universities for ongoing support. We thank the VIP lab of the Palaeosciences Centre and the Microfocus X-ray CT facility of the Palaeosciences Centre at Wits; and for funding these facilities, we thank the University of the Witwatersrand Office of Research and the National Research Foundation Strategic Research Infrastructure Grant and AOP funding programs. For access to comparative specimens, we thank E. Mbua, P. Kiura, V. Iminjili, and the National Museums of Kenya; A. Kwekason, P. Msemwa, and the Tanzania Commission for Science and Technology; B. Billings, Fossil Primate Access Advisory and the School of Anatomical Sciences at the University of the Witwatersrand; S. Potze, L. C. Kgasi, and the Ditsong Museum; Y. Haile-Selassie, L. Jellema, and the Cleveland Museum of Natural History; J. Chupasko and the Harvard Museum of Comparative Zoology; D. Pilbeam, M. Morgan, O. Herschensohn, J. Rousseau, and the Harvard Museum of Archeology and Ethnology; E. Westwig and the American Museum of Natural History; M. Wolpoff and the Department of Anthropology at the University of Michigan; and O. Lovejoy (Kent State University). For technical and material support, we thank Duke University and the University of Zurich 2009 and 2010 Field Schools. Numerous individuals have been involved in the ongoing preparation and excavation of these fossils, including C. Dube, C. Kemp, M. Kgasi, M. Languza, J. Malaza, G. Mokoma, P. Mukanela, T. Nemvundi, M. Ngcamphalala, S. Jirah, S. Tshabalala, and C. Yates. Other individuals who have given significant support to this project include B. de Klerk, W. Lawrence, C. Steininger, B. Kuhn, L. Pollarolo, J. Kretzen, D. Conforti, J. McCaffery, C. Dlamini, H. Visser, R. McCrae-Samuel, B. Nkosi, B. Louw, L. Backwell, F. Thackeray, and M. Peltier. J. Smilg facilitated medical CT scanning of the specimens. The *Au. sediba* specimens are archived at the Evolutionary Studies Institute at the University of the Witwatersrand. All data used in this study are available upon request, including access to the original specimens, by bona fide scientists.

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20 November 2012; accepted 13 March 2013
[10.1126/science.1232999](https://doi.org/10.1126/science.1232999)

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