- 1 Functional Anatomy, Biomechanical Performance Capabilities and Potential Niche of
- 2 StW 573: an *Australopithecus* Skeleton (circa 3.67 Ma) From Sterkfontein Member 2,
- 3 and its significance for The Last Common Ancestor of the African Apes and for Hominin
- 4 Origins
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¹ There is no universal agreement as to whether the separation of humans and their kin from great apes and their kin should be recognised at Family level (Hominidae) or Tribe level (Hominini) or Sub-Tribe level (Hominina). Contributors to this volume have agreed, while choosing their own term, to recognize that disagreement by inserting this footnote.

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49	
50	Abstract (300 words)
51	
52	StW 573, from Sterkfontein Member 2, dated ca 3.67 Ma, is by far the most complete
53	skeleton of an australopith to date. Joint morphology is in many cases closely matched in
54	available elements of Australopithecus anamensis (eg. proximal and distal tibial and humeral
55	joint-surfaces) and there are also close similarities to features of the scapula, in particular, of
56	KSD-VP-1/1 A. afarensis from Woranso-Mille. The closest similarities are, however, to the
57	partial skeleton of StW 431 from Sterkfontein Member 4. When considered together, both
58	StW 573 and StW 431 express an hip joint morphology quite distinct from that of A.
59	africanus Sts14, and a proximal femur of a presumed A. africanus from Jacovec Cavern at
60	Sterkfontein, StW 598. This, and other evidence presented herein, suggests there are two

61 pelvic girdle morphs at Sterkfontein, supporting Clarke (2013) in his recognition of a second

species, A. prometheus, containing StW 573 and StW 431. StW 573 is the first hominid 62 skeleton where limb proportions are known unequivocally. It demonstrates that some early 63 hominins, at the time of formation of the Laetoli footprints (3.6 Ma), were large-bodied. with 64 hindlimbs longer than forelimbs. Modelling studies on extant primates indicate that the 65 intermembral index (IMI) of StW 573, low for a non-human great ape, would have 66 67 substantially enhanced economy of bipedal walking over medium-to-long distances, but that it was still too high for effective walking while load-carrying. It would, however, have 68 somewhat reduced the economy of horizontal climbing, but made Gorilla-like embracing of 69 70 large tree-trunks less possible. Consideration of both ethnographic evidence from modern indigenous arboreal foragers and modern degeneracy theory cautions against prescriptive 71 interpretations of hand- and foot-function, by confirming that both human-like upright 72 bipedalism and functional capabilities of the hand and foot can be effective in short-distance 73 arboreal locomotion. 74

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76 1. Introduction

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While it is now largely accepted that there was no phase of terrestrial knucklewalking in 78 79 hominin evolution (see eg., Dainton and Macho, 1999; Dainton, 2001; Clarke, 2002; Kivell and Schmitt, 2009), and that australopiths show adaptations to both terrestrial bipedalism and 80 81 arboreal locomotion, there is still no firm consensus on whether the 'arboreal' features of australopith postcrania would have been the subject of positive selection or were selectively 82 83 neutral anachronisms (Ward 2002, 2013). The view that the two activities must be substantially mechanically incompatible is still current (see eg., Kappelman et al., 2016). The 84 two alternative paradigms date from extended debates (see eg., Latimer, 1991 versus Stern 85 and Susman, 1991) concerning the significance of the AL-288-1 'Lucy' skeleton of 86 Australopithecus afarensis in 1974. Although some one-third complete, this partial skeleton, 87 and other more recently discovered partial australopith skeletons (eg., the Woranso-Mille 88 89 Australopithecus afarensis skeleton, KSD-VP-1/1 ca. 3.6 Ma [Haile-Selassie et al. 2010]) and the Malapa A. sediba skeletons, MH-1 and MH-2, ca. 1.977 Ma [Berger, 2013]) are too 90 incomplete to provide reliable upper and lower limb lengths, a crucial variable in assessing 91 terrestrial and arboreal locomotor performance capabilities, and lack other auxiliary features 92 which might offer a clear signal of the existence of selection for arboreal performance 93 capabilities. 94

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96 It has become increasingly clear, since the discovery of the foot bones of StW 573 ('Little Foot') some 23 years ago (Clarke and Tobias, 1995), that the continued painstaking 97 freeing of Little Foot's fragile bones from their matrix of hard breccia might for the first time 98 provide unequivocal data on australopith postcranial anatomy. Some missing small skeletal 99 100 elements may be recovered with further excavation, and some (such as the distal foot) were destroyed by lime mining a very long time ago, but at least 90% of the skeleton, complete 101 enough for unequivocal knowledge of limb lengths, and with remarkably good preservation 102 of joint surfaces and other vital detail has now been excavated and prepared (Clarke 2018, 103 104 Figure 26). This completeness is unmatched until Homo ergaster KNM-WT 15000, at 1.5 Ma. After much debate (see Bruxelles et al. 2018, submitted), the age of this specimen is now 105 confidently set at ca 3.67 Ma, very close to the age of the Laetoli footprint trails (Leakey and 106 Hay, 1979), which were previously our best source of information on the locomotor 107 capabilities of australopiths. Now, StW 573 for the first time offers unequivocal information 108 on limb proportions of forelimb and hindlimb. 109

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Here we review aspects of contributions on StW 573 which bear on its potential niche 111 as an individual. These aspects include the geological and palaeoenvironmental context 112 (Bruxelles et al. 2018, submitted), its craniodental anatomy (Clarke and Kuman, 2018, 113 submitted), the endocast (Beaudet et al., 2018a, in press), the inner ear (Beaudet et al. 2018b, 114 submitted), the scapula and clavicle (Carlson et al., 2018, in prep.), the hand (Jashashvili et 115 al., 2018 in prep.) and the foot (Deloison, 2004). Heaton et al. (2018, submitted) focus on 116 statistical/morphometric descriptions of the longbones of StW 573 and comparisons with 117 118 other species, and this paper is referred to where appropriate to avoid duplication. However, the present paper is quite distinct in its focus on: a) comparisons of those aspects of joint 119 120 shape which in the literature are generally regarded as informative concerning locomotor function, and b) reconstruction of StW 573's potential ecological niche by proxy 121 122 experiments and extant in-silico modelling. We adopt the hypothesis-based approach of Wainwright (1991) to species ecomorphology, which focuses on individual performance (and 123 in this respect differs considerably from Bock and Von Wahlert's (eg. 1965) previous 124 formulation which includes form, function and biological role but not performance, and is not 125 therefore suited to experimental testing). This paper, and Heaton et al. (2018, submitted) are 126 127 thus distinct in aims and methods, and while they refer to each other where appropriate and 128 may indeed to be read in tandem as complementary/companion papers, they can equally

stand, and can be assessed, quite independently. In addition, we pay close attention to the
significance of modern ecological dynamics theory, particularly plasticity (see eg. Neufuss et
al. 2014); and neurobiological degeneracy (see eg. Edelman and Gally, 2001), for the
cheiridia (hands and feet) in particular, which suggest that circumspection needs to be applied
to functional interpretation of foot (see eg. Deloison, 2004) and hand morphology.

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Further, euhominoids (otherwise known as crown hominoids, so excluding eg. 135 Proconsulidae, Pliopithecidae etc) display high levels of plasticity in muscle architecture: we 136 have already noted that while DeSilva (2009) asserted that humans could not achieve the 137 required dorsiflexion for chimpanzee-like vertical climbing, Venkataraman et al. (2013) 138 showed that (presumably developmental) fibre-length plasticity enables some forest hunter-139 gatherers to do so, while neighbouring non-climbing populations cannot. Further, Neufuss et 140 al. (2014) showed that while lemurs, like all primarily pronograde mammals studied to date, 141 exhibit a dichotomy in axial musculature between deep slow contracting local stabilizer 142 143 muscles and superficial fast contracting global mobilizers and stabilizers, hominoids, as previously shown for *Homo*, show no regionalization. Thus, it appears that hominoids have 144 145 been under selective pressure to develop and sustain high functional versatility of the axial musculature, reflecting a wide range of mechanical demands on the trunk in orthogrady. 146 147 Neufuss et al. (2014). Using this analytical framework, focusing throughout on StW 573's individual locomotor performance capabilities, we hope to advance understanding of her 148 149 biomechanical interaction with her environment and potential niche: her ecomorphology 150 sensu Wainwright (1991). 151

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154 Background

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2.1. The Sterkfontein Formation

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The Sterkfontein Formation is the world's longest succession documenting the evolution of hominins, fauna, landscape and ecology. Member 5 holds *Homo habilis* and *Homo ergaster*, *Paranthropus*, and evidence for over half a million years' evolution of early stone tool technologies. Member 4, dating to between at least 2.8 Ma. and 2.15 Ma, is the

richest Australopithecus-bearing deposit in the world and contains two species of 162 Australopithecus, together with diverse and extensive faunal evidence. Four metres below, 163 under the yet unexplored and extensive Member 3, which potentially documents a long 164 period of evolution of Australopithecus, lies Member 2, which has yielded the world's most 165 complete Australopithecus skeleton, StW 573, the only hominin found in this deposit. 166 Member 2 formed around 3.67 Ma (Granger et al. 2015) based on isochron burial dating 167 using cosmogenic aluminium-26 and beryllium-10. The detailed descriptions of the Member 168 2 sedimentary units by Bruxelles et al. (2014, 2018 submitted) confirm the 3.67 Ma age of 169 170 the skeleton published by Granger et al. (2015). Pickering and colleagues' recent (2018) repetition of a 2.8 Ma date fails to take into account, and fails to cite, Bruxelles and 171 colleagues' (2014) demonstration that the flowstones are intrusive and dates for StW 573 172 based on them thus invalid. 173 174 Importantly, taphonomic evidence (Clarke 2018 submitted; Bruxelles et al. 2018 175 submitted) indicates that StW 573 died and was fossilized (below the ecological context in 176 which she lived), from a fall into a steep cave shaft leading to an underground cavern. The 177 skeleton is associated, in Member 2, with fauna dominated by cercopithecoids and carnivores 178 179 (see section 2.4). 180 181 2.2 The StW 573 partial skeleton 182 183 The skeletal elements found to date are shown in assembly in Figure 26 of Clarke 184 185 (2018, submitted). The skeleton offers, for the first time in one individual Australopithecus, complete (if deformed) skull and mandible, many vertebrae and ribs, a crushed pelvis and 186 ischiopubic ramus, femora (broken but with overlapping morphology allowing confident 187 length reconstruction), one intact and one slightly damaged but measurable tibia, partial left 188 and right fibulae which overlap sufficiently to be sure of length and morphology, a partial 189 foot (representing primarily the medial column) foot, two scapulae (one articulated with the 190 upper limb), both claviculae (one partial and one complete), both humeri (one partially 191 crushed), both radii and ulnae (one side near-intact and the other crushed and deformed most 192 probably by a badly healed injury in-vivo), and finally one partial and one virtually complete 193 hand (missing only one distal phalanx). 194 195

196 Their taphonomy and condition are discussed in detail in Clarke (2018, submitted) and the stratigraphic context in Bruxelles et al. (2018, submitted). StW 573's pelvis was 197 substantially flattened post-mortem, but preservation of its margins is good enough to 198 identify an obtuse greater sciatic notch angle (Figure 1 top) and hence suggest female sex. In 199 200 contrast Figure 1 bottom shows the original reconstruction of the pelvis of StW 431 from Member 4, which appears closely similar but is indicative of a male from the acuteness of the 201 greater sciatic notch. Lipping of the margins of the vertebral bodies of StW 573 (Figure 1, 202 top) and heavy toothwear (Clarke et al., 2018, submitted), indicate that she was an old 203 204 individual. StW 573 would have been some 130 cm in stature (RJC pers.comm. to RHC), which is some 10 cm. less than the average for modern Bolivian women, the world's shortest 205 female population. By contrast, the stature of AL-288-1 would have been some 107 cm 206 (Jungers, 1988). The considerable difference in stature is in accord with conclusions from the 207 dimensions of the penecontemporaneous Laetoli footprints (both Deloison [1993, pp. 624-208 629], for Laetoli G and Masao et al. [2016] for the more extensive Laetoli S) that there was a 209 large range in stature in early hominins. The slightly younger KSD-VP-1/1 partial skeleton 210 confirms this conclusion for A. afarensis, to which it is referred. 211

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2.3. Environmental Context

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215 While based on the bovids, Vrba (1975) suggested a medium density woodland with a substantial open component, and based on the overall community of mammals, Reed (1997) 216 217 similarly suggested open woodland with bush, However, most of the Sterkfontein Member 2 218 fauna represent cercopithecoid or carnivore taxa which are today habitual climbers, and 219 ancient gravels indicate a large, slow flowing river in the base of the valley (Pickering et al., 220 2004). Similarly, Elton et al. (2016) indicate that the cercopithecoids which are found in 221 Member 2 were probably to some extent ecologically dependent upon trees for foraging, predator avoidance, or both. Thus, Pickering et al., (2004).suggest a paleohabitat of rocky 222 hills covered in brush and scrub, but valley bottoms with riverine forest, swamp and standing 223 water. Such a paleoenvironment might resemble that in today's Odzala-Koukoua National 224 Park, Congo, where grassland, standing water and forest are interspersed (see eg., 225 https://reefandrainforest.co.uk/news-item/trip-report-wildlife-republic-congo. Member 4 226 227 preserves a takin-like (and hence presumably also woodland) bovid *Makapania*, as well as large cercopithecoids (Pickering et al. 2004), which are associated with forest vines requiring 228 large trees (Bamford, 1999), including one today known exclusively from central and 229

230	Western African tropical forest. This evidence suggests that little dessication occurred until
231	Member 5 times. A carbon isotope study of faunal teeth by Luyt and Lee Thorpe (2003) also
232	confirms that a drier, more open environment was only established by 1.7 Ma at Sterkfontein.
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235	2.4. Species affinities
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237	There has been a long debate concerning the number of species within
238	Australopithecus in South Africa, and particularly with relationship to the validity of
239	Australopithecus prometheus. Grine (2013) feels that craniodental and some ancillary
240	paleoenvironmental data are insufficient to justify splitting A. africanus, but Clarke (2013)
241	presents craniodental and also postcranial evidence for species diversity at Sterkfontein. Of
242	particular relevance here, he finds a distinction between A. prometheus, represented by, for
243	example, the partial skeleton of StW 431 and the near-complete StW 573 skeleton and A.
244	africanus, represented by Sts 14. Most of these discussions have focused on dental and
245	gnathocranial evidence, but recent postcranial discoveries of postcrania have broadened the
246	evidence base considerably, with particular attention now being paid to the pelvis and hip
247	joint, which are crucial to both obstetric and locomotor evolution.
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249	Thus, Figure 2 shows that the ilia of the Member 4 StW 431 and the Member 2 StW
250	573 are closely similar in size and shape, even though the greater sciatic notches indicate that
251	StW 573, with an obtuse notch, is most likely a female, and StW 431, with an acute notch, is
252	apparently a male. By contrast, Figure 3 shows that the ilia of Sts14 and A. afarensis AL-288-
253	1 from their obtuse greater sciatic notches both apparent females – are substantially
254	smaller. While Toussaint et al. (2003), who described StW 431, refer it to A. africanus, co-
255	author Macho prefers attribution to A. prometheus (G.A.M. pers. comm. to RHC).

Importantly, both the unreconstructed pelvis described by Toussaint et al. (2003) and the

Kibii and Clarke (2003) reconstruction show that the acetabular margin is well preserved, and
acetabular size well defined. Toussaint et al. (2003) note: 'The acetabulum is clearly large: to

judge by the preserved part, its vertical diameter would exceed 42 mm compared with 29.2
mm in Sts 14' (page 219).

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Acetabular size is also large in StW 573, probably over 36 mm (Figure 4), although the acetabulum has been compressed by taphonomic events. Figures 5 and 6 show that not

only is the acetabulum large, but the StW 573 femoral head is a close match for the 264 acetabulum of StW 431, while the head of the A. africanus proximal femur from Jacovec 265 Cavern StW 598 is markedly smaller than the acetabulum of StW 431 or StW 573. The 266 Jacovec Cavern proximal femur (StW 598) is, however, a good match for the acetabulum of 267 Sts 14 (Figure 7 top left). That this reflects more than allometry is shown by the fact that the 268 269 femur of A. afarensis also has a small head but lacks an obviously long femoral neck. We predict that other isolated material referable to A. africanus will similarly be found to have a 270 long femoral neck. The StW 367 femur from Member 4 Sterkfontein shows a remarkable 271 272 similarity to that of Jacovec StW 598 (Figure 7, top right). Thus both the small-bodied, long 273 femoral neck/small femoral head morph (eg. Jacovec, StW 367) and the large-bodied, short femoral neck but large femoral head large hip joint morph (StW 431, StW 573) were present 274 275 in both Member 2 and Member 4 times. The StW 573 femur resembles those of both humans 276 and KNM WT 15000 (Figure 7, bottom) (and see Heaton et al. 2018, submitted, for 277 morphometric detail). A large femoral head is commonly, and reasonably, associated with large forces operating across the hip joint and may be expected to correlate with body size. 278 279 Femoral neck length, however, is likely related to the moment arm of the hip abductors (see eg. McHenry, 1975), but in what way, and with what iliac geometries, remains to be tested. It 280 281 should be noted that a long femoral neck can prima facie be assumed to increase the risk of 282 femoral neck fracture during instability events or falls, as it will increase the moment arm about the femoral neck from the impact or instability site. Stern (2000) is cited by Toussaint 283 et al. (2003) in reference to the possibility that A. afarensis and A. africanus may have had a 284 less effective abduction capacity in gluteus medius 'thus compromising stabilization during 285 walking' (page 222). This could readily, and will be, tested in silico. 286

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Fornai et al. (2018) recently reported that StW 431 also differs markedly from Sts 14 288 in sacral shape, and, like ourselves, they suggest that functional morphs exist in South 289 African Australopithecus. In this respect, it is interesting that Toussaint et al. (2003) noted 290 very different body mass estimates for StW 431-- 42.5 kg using a hominoid RMA regression 291 292 line, and 41.1 kg using a human regression line, but only 33.4 kg and 22.6 kg respectively on the basis of the lumbosacral region. There are clearly major biomechanical distinctions in the 293 lumbosacral and hip regions, key for the effectiveness of upright walking, between 294 universally recognised A. africanus (eg., Sts 14) and both StW 573 and StW 431. Not all of 295 them can be put down to a simple relationship to body size – although body size itself is a 296 major difference. Since a palaeospecies is identified by morphological (and one would hope, 297

298 functional) distance, we argue that the balance of evidence is now strongly in favor of broad 299 recognition of A. prometheus as a species distinct from A. africanus. 300 301 302 2.5 Cranial and Dental Anatomy and Diet 303 304 An overall cranial shape similarity is evident with the Bouri Hata hominin, ca. 2.5 305 Ma, (Asfaw et al. 1999), but some aspects of cranial morphology suggest to Clarke and 306 307 Kuman (2018, submitted) that an ancestral relationship of StW 573 to Paranthropus may be 308 possible. Beaudet et al. (2018a, in press) conclude from the remarkably well preserved endocast that the brain was small (perhaps surprisingly so) and undistinguished from that of 309 310 other non-human great apes (NHGAs). 311 Although it will take some time for the mandible to be detached safely from the cranium, microCT scanning has revealed that wear distribution as well as dental arcade shape 312 313 resemble those of Kanapoi A. anamensis, 4.17-4.12 Ma., where Ward et al. (2001, p. 351) found that the 'teeth exhibit a distinctive pattern of wear. Evident in older individuals, the 314 anterior teeth are worn very heavily, much more so than the molars and premolars.' Ward et 315 al. (2001) cite evidence that A. anamensis was taking a tough C4 diet, which might suggest 316 open environments. But faunal analysis suggested to Reed (1997) that Kanapoi 317 paleoenvironments at the time of A. anamensis were closed woodland. However, 318 Behrensmeyer and Reed (2013) note that other evidence, including stable isotopes, possibly 319 320 non-arboreal monkeys and micromammals, and characteristics of paleosols, suggest that open habitats also existed. Similarly, Cerling et al. (2013) found that (δ^{13} C) stable isotopes in 321 dental enamel of A. anamensis suggest a C3-dominated diet (leaves and fruits from trees and 322 shrubs, etc.). Further, comparative evidence from extant colobines (Koyabu and Endo, 2010) 323 324 indicates that similar wear distribution may result from consumption of tough-skinned arboreal fruit. Of course consumption of tough-coated arboreal fruit and consumption of 325 326 tough-coated terrestrial resources (such as corms and tubers) are not mutually exclusive. The

Woranso-Mille hominin KSD-VP-1/1, now dated to some 3.6 Ma (Haile-Selassie, 2016) and

- attributed to *A. afarensis*, also appears from faunal evidence to have occupied a primarily
- wooded environment (Su, 2016), with browsers dominant and grazers a relatively small
- component, but with some aquatic species, such as crocodiles and an otter, *Torolutra*,

suggesting that the locality samples a riverbank community. $\delta 13C$ determinations from A.

afarensis at Woranso-Mille suggest a balance of C3 and C4 items in diet (Wynn et al., 2013;

Levin et al., 2015), but microwear (Ungar et al. 2010) closely resembles that in *A*.

anamensis.

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337 **3. Functional Interpretation**

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As noted above, morphometric and general anatomical descriptions of StW 573 long bones 339 are provided by Heaton et al. (2018, submitted); of the scapula by Carlson et al. (2018, in 340 prep.), of the hand by Jashashvili et al. (2018, in prep.), and the foot by Deloison (2004). 341 Detailed descriptions and morphometrics should be sought therein, as here we restrict our 342 attention to the significance of the postcranial anatomy of StW 573 for locomotor ecology of 343 early hominins. We focus our comparative attention primarily on A. anamensis from Kanapoi 344 and KSD-VP-1/1, as they bracket StW 573 in time and are of similar size. StW 431 is 345 younger than StW 573, but we regard this specimen as conspecific. Some comparisons will 346 347 be made to Sts 14 and AL-288-1, but these specimens are considerably smaller and they are more likely to be adaptively different, quite possibly being (in the case of Sts 14 at least) 348 349 more arboreal. We do not refer extensively to the considerably later A. sediba. As noted by Lovejoy et al. (2016), its forelimb seems curiously derived towards some kind of suspensory 350 351 locomotion and/or feeding, and its hindlimbs are not reliably reconstructed, the lower limb length having been assumed by Berger (eg. 2013) to equal that of proximal and distal 352 353 fragments plus the length of the empty matrix between them in situ, despite the likelihood 354 that taphonomic events would affect such an indirect length estimate. We make broad comparisons to the well preserved skeleton of Homo ergaster, KNM WT 15000 (Walker and 355 356 Leakey, 1993), ca 1.5 Mya., as appropriate.

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3.1. Thorax and Pectoral Girdle

The ribs and vertebral column are currently under study by our team but it appears that the thoracic inlet is narrow, unlike the penecontemporaneous KSD-VP-l/1. This does not support the generalization of Lovejoy *et al.* (2016) from KSD-VP-l/1 that early hominins had abandoned the superiorly narrow ribcage typical of NHGAs. On the other hand, the clavicles (Figure 9 top) are broadly humanlike in form, and indeed remarkably long, very similar to

those in the much taller KNM WT 15000 Homo ergaster. The right clavicle is complete 364 (Figure 9 top). Like that of KNM WT 15000 (Figure 9 bottom), it is delicate, with a clear S 365 shape very similar to that exhibited by humans. The strong sigmoid curvatures would 366 increase moment arm for potential stabilizers of the shoulder girdle against the humerus, such 367 as the clavicular head of the pectoralis major, the deltoid, and pectoralis minor. The most 368 remarkable feature, however, is the length: 14 cm, in an early human ancestor estimated to be 369 370 ca. 130 cm in stature (RJC pers. comm. to RHC). This clavicular length equals typical means for adult humans worldwide (Trinkaus et al., 2014) and is in striking contrast with the short 371 372 clavicle of A. sediba MH2 as reported by Schmid et al. (2013).

373 Given the likely close relationship of the male StW 431 to StW 573, we virtually bisected the reconstructed articulated os innominatum and sacrum through the sacral midline 374 375 and mirrored it. The bi-iliac width of the reconstructed StW 431 pelvis (Fig. 1, bottom) was thus estimated at 30 cm. Given the dimensional similarities between the StW 573a and StW 376 377 431 pelves, the bi-iliac breadth of StW 573 cannot have been much less than the 30 cm. biiliac distance in StW 431, which compares to mean values in modern human females of 378 379 around 28 cm. (see eg. Simpson et al., 2008). Since StW 573's clavicle was 14 cm long, and 380 assuming some 3 cm inter-clavicular distance (we lack a sternum), her bi-acromial distance 381 would have been some 28-30 cm, very similar to the likely bi-iliac breadth, suggesting that 382 the trunk was more or less of equal width superiorly and inferiorly, unlike the ribcage. This mismatch between a narrow thoracic inlet and broad shoulders suggests the latter was the 383 subject of active selection for large moments at the glenohumeral joint, and hence powerful 384 climbing. A preliminary canonical variates plot of scapular geometry by Carlson et al. (2018, 385 in prep.) based on MicroCT and virtual reconstruction of the shattered scapular blade shows 386 that StW 573 occupies a position very close to MH2 A. sediba, but also close to KSD-VP-387 1/1, Gorilla and Pongo. However, plots for Pan and Homo (particularly KNM-WT 15000) 388 lie quite distant from StW 573, at the left and right extremes of the plot. The glenoid fossa is 389 390 certainly more cranially oriented than in Homo. Either way, the geometry of the pectoral 391 girdle of A. prometheus does not seem to resemble the 'shrugged' girdle proposed by 392 Churchill et al. (2013) for A. sediba. Weak expression of the mastoid process on the skull of StW 573 (Figure 10) indicates that the sternocleidomastoid was by no means as powerful as 393 394 would be expected with such a 'shrugged' posture. Indeed, the distinction between the short clavicle of A. sediba (1.97 Ma) and the long clavicle of the much earlier A. prometheus (3.67 395 396 Ma) suggests that any elevated pectoral girdle posture in A. sediba is derived, not ancestral as

claimed by Churchill et al. (2013). Following Rein et al. (2017), we must consider whether
suspensory performance was selected for in *A. sediba*, possibly in connection with postural
feeding adaptations.

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401 *3.2 Arm*

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The right humerus is crushed but intact. It is articulated proximally with the scapula 403 and distally with the radius and ulna. The head of the detached left humerus is crushed and so 404 405 the size of the deltoid tuberosity cannot be assessed. Muscle markings are moderately strong, particularly the intact brachioradialis crest (Figure 10 top), which appears substantially larger 406 than the damaged crest in KSD-VP-1/1b figured by Lovejoy et al. (2016). This implies more 407 power in pronation in StW 573 (which hypothesis again can be tested in silico). The distal 408 409 humeral condyles (Figure 10 bottom) appear very similar in form to those of the Kanapoi A. anamensis KNM-KP 271, figured by Hill and Ward (2018) in having, for example, a more 410 411 salient lateral margin for the trochlear articulation than KNM WT 15000 (Figure 11). This feature might imply less axial 'rocking' of the ulna than occurs in our genus, but as Lovejoy 412 413 and colleagues (2016) note, these distinctions are not so major as to necessarily imply active 414 selection. And, as Hill and Ward (1988) note, distal humeral morphology is very variable in humans. Further, Hill and Ward (1988) comment that the Kanapoi distal humerus shows a 415 clear fracture and needs to be considered with caution. 416

417

With regard to the ulna, the shape of the StW 573 trochlear notch agrees more closely 418 with the human ulna figured in Lovejoy et al. (2016) than with either KSD-VP-1/1 or AL-419 288-1 in its somewhat less anterior orientation. Following those authors, we refrain from 420 functional interpretation at this moment. Shaft curvature appears more marked than that 421 422 figured by Lovejoy et al. (2016) for KSD-VP-1/1, but it appears from Drapeau et al. (2005) that curvature is variable in early hominins. There is no radius for KSD-VP-1/1, but that of 423 the Kanapoi A. anamensis, as figured by Ward et al. (2001), is both similar in morphology 424 425 and near-identical in length to that of StW 573. Retention of such a long radius (see section 3.5, *Limb proportions*), especially in combination with a relatively powerful brachioradialis, 426 implies power in flexed/pronated elbow postures, most likely employed during climbing. 427 428

429 *3.3. Pelvic girdle*

430 The os innominatum of both StW 573 and StW 431 corresponds broadly with the form shown by Lovejoy et al. (2016, Fig 8.21) for A. afarensis, with both a greater sciatic 431 432 notch and anterior inferior iliac spine evident, although the latter has sheared off in StW 573. We need not refer further to the pelvic girdle until the crushed pelvis of StW 573 has been 433 434 restored by retrodeformation. Hence most information is drawn from StW 431 (see Toussaint et al., 2002 and Kibii and Clarke, 2003). However, we should note that as Kozma et al. 435 (2018, p. 1) pithily conclude from a study of hip extensor mechanicsm, 'Ardipithecus was 436 capable of nearly human-like hip extension during bipedal walking, but retained the capacity 437 438 for powerful, ape-like hip extension during vertical climbing. Hip extension capability was essentially human-like in Australopithecus afarensis and Australopithecus africanus, 439 suggesting an economical walking gait but reduced mechanical advantage for powered hip 440 extension during climbing.' Contra Lovejoy et al. (2016) who unequivocally attribute a short 441 ischium in *Homo* to running, Kozma et al. (2018) demonstrate that a short ischium greatly 442 enhances distance travelled for energy consumed in walking. But it is worth noting that 443 musculoskeletal modelling by some of us (Goh et al., 2017) showed that in terms of joint 444 moments and torques exerted by all major lower limb extrinsic muscles, the ability of gorillas 445 446 to walk bipedally is not limited by their adaptations for quadrupedalism and vertical 447 climbing.

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- 449

450 *3.4 Femur, Tibia, Hip, Knee and Ankle*

451

We have noted that the femoral head of StW 573 is large, and the femoral neck is 452 453 short compared to A. africanus sensu stricto (e.g., the proximal femur from Jacovec Cavern 454 StW 598 [Partridge et al., 2003]) and A. afarensis AL-288-1. In that respect it resembles KNM-WT 15000 more closely. Unfortunately, there is as yet no proximal femur for the more 455 size and age-matched A. afarensis KSD-VP-1/1. The left distal femur of A. afarensis KSD-456 VP-1/1 is poorly preserved, especially the medial condyle, but Lovejoy et al. (2016) report 457 that the (restored) lateral condyle is 'elliptical', and like StW 573, the patellar groove is deep 458 and shows a high lateral wall for patellar retention, as noted by Heaton et al. (2018, 459 submitted). 460

461

However, the lateral femoral condyle of StW 573 (Figure 12) is not only posteriorly
'elliptical' (to use Lovejoy and colleagues' [2016] term), but more specifically like humans,

has a relatively rounded posterior/dorsal section and flat anterior/ventral section. Again like

humans, the medial femoral condyle is more evenly rounded dorsoventrally (Figure 13). The

466 knee of KSD-VP-1/1 does, as Lovejoy et al. (2016) state, appear to show a valgus angle (see

their Figures 8.6 and 8.6). But it is, like that of StW 573 (Figure 14) more weakly marked

than in KNM-WT 15000 and particularly than in AL-288-1 (Stern and Susman 1983), where,

- taken at face value, the angle probably reaches an extreme among hominins.
- 470

471 There is detailed evidence of the morphology of the proximal surface of the A. anamensis tibia from Kanapoi KNM-KP 29285A, 4.16 Ma (reviewed in Ward et al., 2001), 472 which is shown in Figure 15 as a visualization of an stl file (open source, from: 473 africanfossils.org. XYZ dimensions 68.00; 103.30; 60.66 mm.). Ligamentous and muscular 474 attachments are detailed, but although preservation of StW 573 and KNM WT 15000 is 475 excellent, these are not identifiable with any confidence in either specimen (Figure 16 top and 476 bottom). The tibia of KSD-VP-1/1 is heavily damaged throughout and the proximal surface 477 carries little information. From Figures 17, 18 and 19, it is clear that KNM-KP 29285A, StW 478 479 573 and KNM WT 15000 all have long, concave condyles on the medial side, and short, less 480 concave condyles on the lateral side, which in KNM-WT 15000 and StW 573 are matched by a long rounded section on the medial femoral condyle but an anteriorly flatter lateral condyle. 481 482 This is the bony basis of the 'locking' or 'screw-home'mechanism of the knee (see eg. Dye, 1987 and Lovejoy, 2007). The condyles and cruciate ligaments form a four-bar linkage. In 483 484 knee extension, because of the flatter condylar morphology of the ventral part of the lateral condyles, they cease sagittal rotation motion before the medial condyle, and rollback occurs, 485 486 compressing the lateral meniscus and further immobilizing the lateral condyle so that a 487 passive coronal rotation results, spiralizing fibres in the cruciate ligaments and stabilizing the 488 knee. This allows standing with minimal expenditure of muscular energy for balance and 489 signifies that early hominins from 4.16 Ma onwards (including both A. anamensis and A. prometheus) were able to stand upright with enhanced efficiency. ARA VP1/701 490 Ardipithecus ramidus lacks most of the femur, and curiously, the nearly complete tibia is 491 largely unreported (see eg. White et al., 2009), so we cannot assess whether Ar. ramidus had 492 this important mechanism, despite Lovejoy's reference to his own (2007) paper discussing 493 the so-called 'screw-home' mechanism reviewed above. But as might be expected the 494 495 associated distal femoral condyle asymmetry is evident in the morphology of the AL-288-1 distal femur, 3.4 Ma (Figure 20) (and see Stern and Susman, 1983 and Lovejoy, 2007). 496

In upright arboreal bipedalism, some of us have shown experimentally (Johannsen et
al., 2017) that 'light touch' with the fingers on supports between shoulder and waist height
significantly enhances balance on unstable supports and reduces thigh muscle activity
required to counteract perturbation by some 30%. *Ar. ramidus* could have used this
mechanism in upright bipedal walking in the trees, and thus it could have been an effective
upright arboreal biped. The same applies to *Pierolapithecus* (12.5-13 Ma, Moyà-Solà et al.,
2004).

504

Lovejoy et al. (2016) draw attention to the short radius of curvature in the talar joint 505 surface of the distal tibia KSD-VP-1/1, , versus the flatter talar joint surface in NHGAs 506 507 (DeSilva, 2009). Figure 21 shows that the radius of curvature is as short in StW 573 as it is in KNM-WT 15000, and a similarly short radius of curvature can be seen in Figure 22 (an stl 508 model of Kanapoi distal right tibia KNMKP 29285, downloaded from [open source] www. 509 africanfossils.org, Dimensions: x=40.16; y=97.82; z=40.50 mm). Ward et al. (2001) note 510 that the maximum concavity of that Kanapoi talar joint surface/plafond is 5 mm. In StW 573 511 it is ca 4.5 mm, and in KNM-WT 15000 (depending on side) it is also ca 4.5 mm. In each 512 513 case the shape of the talar joint surface is square, rather than rectangular as tends to be the 514 case in NHGAs.

515

516 DeSilva (2009) claimed that the human ankle joint was incapable of dorsiflexion to the extent required for 'chimpanzee-like' vertical climbing, and this view has been widely 517 518 taken on board, particularly by Lovejoy et al. (e.g., 2016). However, Venkataraman et al. (2013a) showed that Twa hunter-gatherers can indeed achieve high ankle dorsiflexion, and 519 520 engage in vertical climbing since they tend to have longer fibers in the gastrocnemius muscle 521 than neighbouring, non-climbing agricultural communities. The latter is an excellent example 522 of the importance of plasticity - the ability to adapt musculoskeletal anatomy during development to enhance function in the realized niche - to all great apes, including humans, 523 to which we shall return later. 524

StW 573, KNM-WT15000 and Kanapoi *A. anamensis* thus appear to have very
similar proximal and distal tibial morphology, which strongly suggests similarity in function.
However, isolated and species-unidentified specimens from Sterkfontein Member 4 often
show rather variable morphology. The Member 4 specimen StW 514 assumed to be *A. africanus* by Berger and Tobias (1996), however, combines an *A. anamensis*-like distal tibial

530 condyle (StW 514b) with a proximal condyle (StW 514a), which Berger and Tobias (1996) claimed had distinctly more convex condyles than A. afarensis. Organ and Ward (2006), 531 however, found no difference in lateral tibial condyle geometry between StW 514a and A. 532 afarensis, and the debate concerning whether, and to what extent, Member 4 australopiths 533 534 developed a wider range of locomotor adaptations continues. The case of the peculiar pectoral girdle adaptations of A. sediba from Malapa, for example (Churchill et al., 2013) is 535 strong evidence that some South African species may have adopted unique modes of postural 536 537 feeding. 538 539 3.5. Limb proportions 540 541 542 Figure 23 shows the long bones of the upper and lower limbs of StW 573 compared. At a likely 130 cm tall (RJC pers. comm. to RHC), she was some 10 cm shorter than the 543 544 average for modern Bolivian women, but some 23 cm taller than AL-288-1 Lucy (106.68 cm 545 according to Jungers, 1988). She was a little shorter than A. afarensis KSD-VP-1/1, by the 546 margin which might be expected in a female. Her left humerus' maximum length is 29 cm; 547 her radius is 24.4 cm long (RJC pers. comm. to RHC), almost identical to the length of the A. anamensis radius from Allia Bay, East Turkana, KNM-ER 20419, which it also resembles 548 closely in its (conservative) morphology. Her ulna was 26.3 cm. long. Her total arm length 549 (humerus plus radius) was 53.4 cm. Her femora would have been 33 cm in length, 28.5 for 550 551 the tibia, giving a total leg length of 61.5 cm (RJC, pers. comm. to RHC and see Heaton et al., 2018 submitted). Thus it is no longer a subject for debate whether some early hominids, 552

living at about the time the Laetoli G and S trails were laid down, had hindlimbs that were as
long or longer than their forelimbs. StW 573 is the first hominin fossil in which this is
unequivocal.

556

These values give an intermembral index of 86.8 (ratio of h + r L to f + t). Other indices are discussed in Heaton *et al.* (2018 submitted). This is outside and above the human range as reported by Schultz (1937) at 64.5-78, but below that of *Gorilla* at 110-125, that of *Pongo* at 135-150.9, and that of *Pan* at 100.4-100.5, but clearly much closer to the human range than that of the other great apes. The range in *Pan* is so narrow compared to all other great apes as to suggest it is under strong selective control, most likely very tight tuning for

563	effectiveness in quadrupedalism (see Isler et al. 2006). Indeed Drapeau and Ward (2007)
564	note that the proportions of the forelimb in <i>Pan</i> are highly derived.
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568	4. Discussion
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570	4.1 Ecomorphology: Testable Hypotheses on Potential Niche
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572	Bock and von Wahlert's (1965) classic paper, 'Adaptation and the Form-Function Complex,'
573	stressed form, function and biological role. It inspired a generation, some to explore
574	biological role by field studies in the natural environment, and others to pursue analyses of
575	the biomechanics of living primates held in captivity. Despite this, it may fairly be said to
576	have had relatively little influence in changing methodology in hominin paleontology, where
577	morphometrics now most often geometric morphometrics continues to dominate research
578	activity, although the introduction of biomechanical modelling techniques such as Finite
579	Elements Stress Analysis and Dynamic Modelling, has been pursued by a (growing)
580	minority. In our view, a newer ecological formulation that is hypothesis- and experiment-
581	driven would be greatly beneficial. This was provided by Wainwright's (1991)
582	'Ecomorphology: Experimental Functional Anatomy for Ecological Problems'. It updates
583	Bock and Von Wahlert (1965) in its focus on performance, and specifically performance of
584	the individual, which is vital because it is the reproductive success of the individual which
585	drives adaptation at population and species levels. Wainwright (1991, p. 680) says:
586	'morphology influences ecology by limiting the ability of the individual to perform key tasks
587	in its daily life. In this scheme the effect of morphological variation on behavioral
588	performance is first tested in laboratory experiments. As the behavioral capability of an
589	individual defines the range of ecological resources that it can potentially make use of (the
590	potential niche), the second step in the scheme involves comparing the potential niche of an
591	individual to actual patterns of resource use (the realized niche)'. Those of us who study
592	fossils can rarely carry out 'laboratory experiments on the effect of morphological variation
593	on behavioural performance' (p. 680), but increasingly, we can do so in silico, using custom-
594	designed-and-written software. This is usually open-source, such as OpenSim
595	(http://opensim.stanford.edu/work/index.html) and co-author Sellers' GaitSym
596	(<u>www.animalsimulation.org</u>). The latter has been specifically written for comparative, not

597 human biomechanics, and for palaeontology. Another approach is experimentation using human proxies: we have cited one such study, Johanssen et al. (2017), which tested the effect 598 of light touch on stabilization of the body on unstable supports in a visual simulation of 599 rainforest environments. Similarly, we have just reported that upper limb lengths were short 600 in StW 573 compared to the NHGAs. This suggests less ability to embrace large supports, 601 and particularly, shorter reach, which we hypothesize to reduce the energetic efficiency of 602 arboreal locomotion. Halsey et al. (2017) measured the impact of variation in morphology 603 604 and locomotor behaviour on the rate of oxygen consumption of 19 elite male parkour athletes 605 as they repeatedly traversed an arboreal-like assault course of 103 m horizontal length. The course consisted of a range of generic gymnasium apparatus such as vaulting horses, raised 606 blocks, high bars, wall bars, and areas filled with loose foam blocks to emulate the range of 607 mechanical conditions present in an arboreal pathway, rather than the exact structure of the 608 forest canopy. Thus, parts of the course incorporated support compliance, irregularity and 609 discontinuity to reflect the conditions experienced during gap crossing between tree crowns, 610 while others were rigid and predicated to reflect the phases between bouts of gap crossing 611 612 when even large-bodied apes may walk into and out of the core of a tree along thick boughs. They found familiarity with the course had a substantial effect on reducing energetic costs, 613 614 but there was no evidence to suggest that the locomotor behavior profile of each individual (or the combination of locomotor behaviors that they selected between first and last trials) 615 616 influenced their ability to attenuate costs. We must therefore, presume more subtle mechanical adjustments are being made to attenuate locomotor challenges. Importantly, 617 618 athletes with longer arm spans and shorter legs were particularly able to find energetic 619 economies. Thus, our hypothesis that shorter reach would reduce the efficiency of arboreal 620 locomotion is confirmed for one hominin at least, namely Homo sapiens. Therefore based on this analogy we conclude that the limb proportions of StW 573 would have reduced her 621 622 energetic efficiency in arboreal climbing.

623

A second hypothesis would then be that her long legs and shorter arms would have increased her distance-specific effectiveness in bipedalism. While we have commenced insilico modelling of StW 573 using sophisticated forwards dynamics modelling under GaitSym, successful fully 3D modelling inevitably takes a great deal of iterative computation, and thus time. However, previous studies of other hominins and of the biomechanical consequences of their body and limb proportions provide strong indications of likely findings. Wang and Crompton (2003, Figure 3), using mass and stature estimates from the 631 literature, found that dimensionless power, mass and stature are closely related, and that humans have arrived at a better combination of these parameters for long distance bipedalism 632 than KNM WT 15000, AL-333 and SK 82. However, as shown by Wang and Crompton 633 (2003, Figure 3) all these fossils occupy a considerably more optimal place on a 3D plot of 634 dimensionless power, mass and stature than for example AL-288-1, OH 62 (a supposed 635 Homo habilis—Johanson et al., 1987, but see Clarke, 2017 for a view contra) and Sts 14. 636 Given her estimated stature (130 cm), StW 573 would occupy a position closer to KNM-WT 637 15000 and AL-333 than to Sts 14, OH 62 and AL-288-1. Thus, our second hypothesis is 638 639 confirmed by analogy: that StW 573's distance-specific effectiveness in bipedalism would be 640 enhanced by her longer legs.

641

On the other hand, following the calculations of Wang et al. (2003), StW 573's intermembral 642 index of 86.8, outside the human range and larger than that of KNM-WT15000, would not 643 have allowed her to hand-carry loads more than the weight of the upper limb without losing 644 swing symmetry. This contrasts with their estimate that KNM-WT 15000 could effectively 645 carry loads of three times the weight of the upper limb while maintaining swing symmetry. 646 Interestingly, chimpanzees proved unable to hand-carry loads at all without losing swing 647 648 symmetry, which is interesting in the light of data showing manuports used by chimpanzees in cracking Panda oleosa nuts in the Taï forest are carried no more than 10-15 m (Profitt et 649 650 al., 2018). Similarly, but using inverse dynamics and shoulder-borne loads, Wang and Crompton (2004) showed that, for the given body proportions, KNM-WT 15000 could carry 651 loads of 10-15% body mass for no greater mechanical cost than AL-288-1 would incur 652 walking upright but unloaded. StW 573 would, we predict, function better in this regard than 653 654 AL-288-1, but by no means as well as KNM-WT 15000. This strongly suggests that her performance capabilities balanced distance-specific terrestrial effectiveness against retention 655 656 of efficiency in arboreal climbing.

657

These hypotheses need to be tested, and currently are being tested for StW 573, using forwards dynamic modelling. Again, we can use this technique to discover what advantage would have been delivered to StW 573 by her short femoral neck, given her substantial pelvic flare. Further, as suggested above, given her more cranially oriented glenoid fossa and scapular form (similar to that of *Gorilla* as well as *A. sediba* and KSD-VP-1/1), but her long clavicles (very unlike *A. sediba*) we can use dynamic modelling of her own unique pectoral girdle and pectoral limb architecture to explore the power that she could exert in moderately 665 elevated glenohumeral postures. Thus we will assess the hypothesis that her large

brachioradialis flange (suggesting a semiflexed/semipronated elbow posture) would

667 maximize flexor power. This would facilitate climbing on narrow diameter treetrunks and

- vines with similar kinematics to that recorded for modern human indigenous arboreal
- 669 foragers, particularly when using hallucal grasping.
- 670
- 4.1. Implications of ecological dynamics for functional capabilities of hands and feet
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Ecological dynamics seeks to explain coordination and control processes in
movement systems during performance of complex multi-articular tasks. This is ner more
obvious than during grasping and stepping where multiple rows of bones each form multiple
joints controlled by many ligaments and tendons (see Seifert et al., 2016). Here, the hands
and feet are interacting directly with the environment and with technology.

The ability of anatomically and morphologically complex organs to adapt efficiently 677 to changes in the environment is driven by the evolutionary mechanism of neurobiological 678 degeneracy. This is the ability of biological elements that are structurally different to perform 679 similar functional outputs (Edelman, 1987). It is quite different to the common engineering 680 681 concept of redundancy, which refers to the duplication or repetition of similar shaped 682 elements to provide alternative functional outputs in times of mechanical failure (Bernstein, 1967). Therefore multiple means of achieving the same or different functions (according to 683 684 ecological context) exist by recruitment of structurally different elements. Neurobiological degeneracy 'is a prominent property of gene networks, neural networks, and evolution itself' 685 686 (Edelman and Gally, 2001, p. 13763).

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688 Further, euhominoids/crown hominoids (i.e., Hominidea excluding eg. Proconsulidae, Pliopithecidae etc.) display high levels of plasticity in muscle architecture. Venkataraman et 689 690 al. (2013) showed that (presumably developmental) fibre-length plasticity enables some human forest hunter-gatherers to dorsiflex the ankle to the extent required for chimpanzee-691 like vertical climbing, while neighbouring non-climbing populations cannot. Further, 692 Neufuss et al. (2014) showed that while lemurs, like all primarily pronograde mammals 693 studied to date, exhibit a dichotomy in axial musculature between deep slow contracting 694 695 local stabilizer muscles and superficial fast contracting global mobilizers and stabilizers, 696 hominoids, as previously shown for *Homo*, show no regionalization. Thus, it appears that 697 hominoids have been under selective pressure to develop and sustain high functional versatility of the axial musculature, reflecting a wide range of mechanical demands on the 698

trunk in orthogrady. Neufuss et al. (2014) suggest that this is a derived characteristic acquired
by early euhominoids. Most likely, this characteristic was acquired by euhominoids such as *Morotopithecus*, or at least *Pierolapithecus*.

702

703 Thus, locomotor flexibility is a characteristic of the euhominoid/crown hominoid clade. But in individuals, degeneracy not only stabilizes under perturbation as in light touch 704 705 (see eg. Johanssen et al. 2017), but helps individuals exhibit adaptability. Multiple alternative recruitment patterns exist in the motor control system, and are variously selected 706 707 by the CNS (central nervous system) in each grasp or step, as the CNS seeks to optimize 708 performance. It results in functional intra-individual movement variability (Seifert et al., 2014). Thus, it is unsurprising that high intra- and inter-subject variability in human foot 709 pressure cannot be characterized reliably by less than 400 step trials (McClymont 2016; 710 McClymont and Crompton, submitted ms.). Such variability is a natural product of a 711 degenerate system so that, for example, even in small samples, peak midfoot pressures 712 overlap in human, bonobo and orang-utan populations (Bates et al., 2013). Further, 713 prehensive capabilities of the human foot need to be assessed in the context of the greater 714 715 abduction of the hallux known for many years to exist in habitual barefoot walkers such as 716 Hoffman's (1905) indigenous forest foragers (and see D'Août et al., 2009). Indigenous human arboreal gatherers such as the Ba'aka, Twa and Batek have the ability to climb small vines 717 718 using a hallucal grasp (see eg. Figure 25), as observed by Kraft et al. (2014), and equally that of Western adults with reduced pollical capabilities or no pollex to substitute skilled hallucal 719 720 grasping. Figure 26 illustrates the refined grasp that can be performed by the hallux of some such individuals. The latter, in particular, is an excellent demonstration of how 721 722 neurobiological degeneracy allows the foot to perform the many fine locomotor skills we 723 tend to associate with the hand. Figure 27 demonstrates that parkour athletes can perform 724 brachiation on an I-beam (here demonstrating the range of plasticity which exists in human finger capabilities, in performing behaviors we normally associate with gibbons and NHGAs, 725

The relative proportions of the thumb and fingers of StW 573 (Figure 28) are modernhuman-like (Clarke, 1999), as is the case with the *A. afarensis* hand from AL 333 and AL 333w, according to Alba et al. (2003). This suggests that modern human-like hand proportions , as well as grasping capacities (Clarke, 1999, 2002) had their origins in arboreal behaviour before they were exploited in more terrestrial hominins for tool-use. Clarke (2002) notes that no stone tools have been found in Member 2, and there is no suggestion that StW 573 made stone tools. On the contrary, Little Foot's hand bears a salient apical ridge on the
trapezium, a feature commonly present, and marked, in living gorillas (Figure 29). This might
have reduced effectiveness of deep, soft opposition (for discussions of prehension see eg.
Marzke et al., 1997, and Tocheri et al., 2008).

Trapezium morphology is highly variable in primates (Napier and Davis, 1959; 736 737 Hellier and Jeffery, 2006), so care must be taken in interpretation, but it is likely that this structure, absent in humans, might help brace the thumb and its ulnar and radial 738 carpometacarpal and metacarpophalangeal collateral ligaments against forced abduction, 739 similar to 'gamekeeper's thumb' which tends to affect skiers who fall on their hand while still 740 grasping their poles, or football (soccer) goalkeepers who fall while holding a football 741 (Glickel, Barron and Eaton, 1999). In gorillas, the apical ridge might therefore stabilize the 742 pollex in abducted pinch grips during climbing, and we suggest that the case would be the 743 744 same in StW 573.

745

746 Available footbones of StW 573 have been discussed in detail by Clarke (1998, 747 2002), Clarke and Tobias (1995) and Deloison (2004). Proportions and general morphology 748 broadly resemble those from Woranso-Mille (Haile-Selassie et al., 2012) and Dikika 749 (DeSilva et al., 2018), and the high functional plasticity of the human hallux discussed above must be taken into account in any discussion of hallucal function. Human feet as a whole are 750 highly plastic and functionally degenerate, and as shown by Venkataraman et al. (2013b) 751 and Kraft et al. (2014), they are perfectly capable of functioning efficiently in climbing as 752 753 well as terrestrial bipedal walking and running, having unquestionably retained a prehensile (if relatively adducted) hallux (see e.g. Figures 25 and 26), contra Holowka and Lieberman 754 755 (2018). The high human death rates from falls from trees of less than 20 m. quoted by 756 Venkataraman et al. (2013b) are a clear indication that, even were plasticity and degeneracy 757 insufficient, selection would certainly favour retention of hallucal prehension in any human population engaging in barefoot climbing (common in human childhood). It is also highly 758 759 pertinent to this discussion that analyses of the Laetoli G1 and G2 footprint trails, both of which were formed by hominins penecontemporaneous with StW 573 and KSD-VP-1/1, 760 761 show that only for very small areas of the foot can external function be statistically 762 distinguished from those made by Holocene human pastoralists and Western humans 763 (McClymont, 2016; McClymont and Crompton submitted ms.). This indicates that the 764 external function of the foot during terrestrial bipedal walking has changed very little since

the time of StW 573. Preliminary studies by Raichlen and Gordon (2017) for the new LaetoliS trails are in agreement with this conclusion.

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- 768

4.2 Significance of StW 573 for Hominin origins and the Last Common Ancestor of African Apes

In summaries of the findings in the 2009 special issue of *Science* on *Ardipithecus* 771 772 ramidus, Lovejoy (p. 74e1) claims 'Ar. ramidus was already well-adapted to bipedality, even though it retained arboreal capabilities. Its postcranial anatomy reveals that locomotion in the 773 chimpanzee/human last common ancestor (hereafter the CLCA) must have retained 774 generalized above-branch quadrupedality, never relying sufficiently on suspension, vertical 775 climbing, or knuckle walking to have elicited any musculoskeletal adaptations to these 776 behaviors.' While we agree strongly with Lovejoy's (2009) view, expressed elsewhere in the 777 778 same paper, that the human/chimpanzee ancestor was not chimpanzee-like, at least in 779 postcranial morphology, we differ with his conclusion that the Pan/Homo LCA must have retained 'generalized above-branch quadrupedality'. *Pan*'s forelimb morphology is highly 780 781 derived (Drapeau and Ward, 2007) and its intermembral index optimized for quadrupedalism (Isler et al., 2006). Why should the LCA not have been a 'well-adapted' arboreal biped as 782 783 some of us (eg. Crompton et al. 2010) have suggested from field data on Pongo and Gorilla locomotion? The work of Johannsen et al. (2017) demonstrates clearly that humans retain 784 785 neural mechanisms for fast response to perturbation in bipedalism on narrow, unstable supports via light touch with the fingers. These would be completely incompatible with a 786 787 hand loaded in quadrupedal posture.

The skeletal similarity of StW 573 to KSD-VP 1/1 and particularly A. anamensis, and 788 789 evidence for a similar diet to the latter -- substantially C3 foods -- suggests that these hominins had a similar potential niche. It further suggests that the contemporaneous Laetoli G 790 791 and S trails were made by a very similar hominin which combined continued, if uniquely 792 hominin, modes of arboreal foraging -- in mesic environments -- with effective terrestrial 793 bipedalism. While Ward et al. (2001) concluded that A. anamensis was very largely terrestrial, they made a point of not ruling out a substantial arboreal component in its ecology. 794 795 The postcranial evidence shows that selection was operating on A. prometheus to retain considerable arboreal competence: from limb proportions, through the long radius shared 796 797 with A. anamensis, (as indicated by the KNM-ER 20419 Sibilot radius from Allia Bay [see

Ward et al. 2001]) to the apical ridge on the trapezium. Indeed the retention of an inner-ear
mechanism suited for motion in a complex, 3D environment demonstrated by Beaudet et al.
(2018b, submitted) is clear endorsement of the interpretation of a substantially arboreal
habitus for *Au. prometheus*. We are thus now able to confirm that the apparent 'arboreal'
features of early hominins were indeed the subject of positive selection, not selectively
neutral anachronisms (see Ward 2002, 2013).

Frequent skeletal similarities of the StW 573 postcranium (e.g., the scapula) to 804 Gorilla gorilla, lacking in Pan, suggest availability of a similar potential niche, but with 805 reduced use, compared to Gorilla, of large tree trunks and increased use of vines and small 806 807 treetrunks, as noted by Venkataraman et al. (2013b) for living human arboreal foragers. Thus we differ also with White et al. (2009, p. 64) in their scenario, which places Gorilla on an 808 'adaptive pedestal' separated from australopiths by the chimpanzees, which suggests 809 unidirectional evolution of hominin locomotion. Pan is biomechanically highly derived. It is 810 811 clear that effective arboreal, as well as terrestrial, foraging, albeit less effective than in 812 NHGAs due to adaptations for increased terrestrial effectiveness, were part of the australopith niche and, given locomotor plasticity and degeneracy, remain part of the potential niche of 813 814 Homo sapiens (Kraft et al., 2014).

815

816 **5.** Conclusions

Following Wainwright's (1991) formulation of ecomorphology, we predict that StW 817 573's potential niche was exploitation of both arboreal and terrestrial resources, facilitated by 818 plasticity and degeneracy. Toothwear and postcranial similarities to A. anamensis suggest a 819 820 similar primarily C3 diet in mesic mixed forest/grassland. This might include fibrous tubers on the ground and at water margins, as well as tough-skinned arboreal fruit. StW 573 was an 821 822 effective arboreal biped and climber which had, however, sacrificed some arboreal 823 effectiveness in favour of enhanced energetic efficiency in walking medium to long distances 824 on the ground. She would not have been as effective when load carrying, unlike Homo ergaster. Her locomotor posture was upright bipedalism, whether on the ground or on 825 826 branches, and she was able to stand upright without much muscular activity because of a 'locking' or 'screw-home' mechanism in the knee which does not seem to have been present 827 in Ar. ramidus. A. anamensis and KSD-VP-1/1 probably shared a similar niche. However, we 828 require new fieldwork on lowland gorilla arboreality to establish how the realized niche of A. 829

prometheus, *A. anamensis* and *Ar. ramidus* in arboreal foraging might have differed from that
of *Gorilla*, accompanied by in-silico testing of locomotor hypotheses concerning early

832 833

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hominin performance capabilities.

835

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1180	Figures
1181	
1182	All images from casts unless otherwise stated
1183	
1184 1185 1186	1 Top: Pelvis of StW 573, Bottom: the Robinson (1972) reconstruction of the pelvis of StW 431 (original photograph)
1180	2 Innominates of (Left) StW 431 (Right) StW 573
1188	
1189 1190	3 Innominates of (Left to Right) StW 431, Sts14 and AL-288-1
1191	4 Innominate of StW 573 showing acetabulum
1192 1193 1194	5 Top, proximal femur of StW 573 mounted in its acetabulum; bottom, mounted in
1195	acetabulum of StW 431 innominate
1196	
1197	6 Top, the proximal femur from Jacovec Cavern StW598 mounted in acetabulum of StW
1198	573; bottom, the proximal femur of StW 573 mounted in acetabulum of the Sts 14 innominate
1199	
1200	7 Figure 7 Top left, the proximal femur from Jacovec Cavern StW598 mounted in the
1201	acetabulum of Sts 14; Top right, the StW367 proximal femur from Member 4; Bottom;
1202	Proximal femora of (left to right) Jacovec StW598, AL-288-1, StW 573 and KNM WT 15000
1203	
1204	8 Top, the right clavicle of StW 573, Bottom, the right clavicle of KNM WT 15000

1205						
1205	9 The mastoid process (arrow) on the skull of StW 573					
1200	5 The mustere process (arrow) on the skull of 5tw 575					
1209 1208 1209	10 Distal humerus of StW 573. Top, ventral view showing brachioradialis crest; bottom, dorsal view showing shape of distal condyles					
1210						
1211	11 Distal humerus of KNM WT 15000, dorsal view					
1212						
1213	12 Lateral distal femoral condyle of StW 573					
1214						
1215	13 Medial distal femoral condyle of StW 573					
1216						
1217	14 Valgus angles of the knee in, left to right: KNM WT 15000, AL-288-1 and StW 573					
1218						
1219	15 Proximal tibial surface of Kanapoi KNM KP 29285A					
1220	(downloaded from open source: www.africanfossils.org,					
1221	XYZ dimensions 68.00; 103.30; 60.66 mm)					
1222						
1223	16 Proximal tibial surface of (Top) StW 573 and (Bottom) KNM WT 15000					
1224						
1225	17 (Top) Frontal, (Bottom left) Lateral, and (Bottom) Medial perspectives of Kanapoi					
1226	KNM KP 29285A (downloaded from open source: www.africanfossils.org, XYZ dimensions					
1227	68.00; 103.30; 60.66 mm)					
1228						
1229 1230	18 (Top) Frontal, (Middle) Lateral and (Bottom) Medial perspectives of proximal tibial condyles of StW 573					
1231						
1232	19 (Top) Frontal, (Middle) Lateral and (Bottom) Medial perspectives of proximal tibial					
1233	condyles of KNM WT 15000					
1234						
1235	20 (Top) Axial, (Middle) Lateral and (Bottom) Medial perspectives of distal femoral					
1236	condyles of AL-288-1					
1237						

1238	21 Radius of curvature of the distal tibial condyle/plafond
1239	Top: StW 573 Bottom: KNM WT 15000
1240	
1241	22 Kanapoi distal right tibia KNM KP 29285 from lateral side, ((downloaded from open
1242	source: <u>www.africanfossils.org</u> ,
1243	Dimensions: x=40.16; y=97.82; z=40.50 mm.)
1244	
1245	23 Long bones of the upper and lower limbs of StW 573
1246	
1247	24 Figure 24 Archival image of an indigenous arboreal forager climbing a thin vine using
1248	flexed elbow postures and hallucal grasp (courtesy of Kirk Endicott)
1249	
1250	
1251	25 An indigenous Batek arboreal forager demonstrating his hallucal grasp for climbing a
1252	small vine (video frame, courtesy of Vivek Venkataraman)
1253	
1254	26 Frames from the official video ('Drawin' on Darwin' - Live drawing created at the Meeting
1255	of the Primate Society of Great Britain Saranjit Birdi Axisweb) of :Top, the artist Saranjit
1256	Birdi drawing with his foot Bottom: closeup to show precision of hallucal grasp (courtesy of Emily Soundary and the artist)
1257	Emily Saunders and the artist)
1258	
1259	
1260	27 Video frame of a parkour athlete brachiating on an I-beam, demonstrating the plasticity of
1261	human finger capabilities (see Halsey et al. 2017, permission of the subject and courtesy
1262	Susannah K.S. Thorpe)
1263	
1264	28 Handbones of StW 573
1265	
1266	28 Apical ridge on the trapezium of: left Gorilla gorilla beringei, right StW 573 (originals)

1267 Figure 1 Top: Pelvis of StW 573, Bottom: the Robinson (1972) reconstruction of the pelvis of

1268 StW 431 (archival photograph)





1278 Figure 2 Innominates of (Left) StW 431 (Right) StW 573



- 1294 Figure 3 Innominates of (Left to Right) StW 431, Sts 14 and AL-288-1



1313 Figure 4 Innominate of StW 573 showing acetabulum



- 1317 Figure 5 Top, proximal femur of StW 573 mounted in its acetabulum; Bottom, mounted
- 1318 in acetabulum of StW 431 innominate



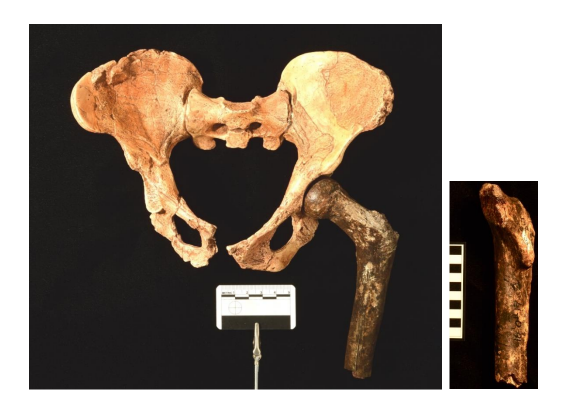


- 1325 Figure 6: Top, the proximal femur from Jacovec Cavern StW 598 mounted in acetabulum
- 1326 of StW 573; Bottom, the proximal femur of StW 573 mounted in acetabulum of the Sts
- 1327 14 innominate





Figure 7: Top left, the proximal femur from Jacovec Cavern StW 598 mounted in the acetabulum of Sts 14; Top right, the StW 367 proximal femur from Member 4; Bottom; Proximal femora of (left to right) Jacovec StW 598, AL-288-1, StW 573 and KNM WT 1337 15000





- 1345 Figure 8 Top: The right clavicle of StW 573 Bottom: The right clavicle of KNM WT 15000





1356	Figure 9: The mastoid	process	(arrow) o	on the	skull	of StW :	573
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- 1366 Figure 10: Distal humerus of StW 573. Top, ventral view showing brachioradialis crest;
- 1367 Bottom, dorsal view showing shape of distal condyles





1373 Figure 11: Distal humerus of KNM WT 15000, dorsal view

1374



1377 Figure 12: Lateral distal femoral condyle of StW 573



1388 Figure 13: Medial distal femoral condyle of StW 573

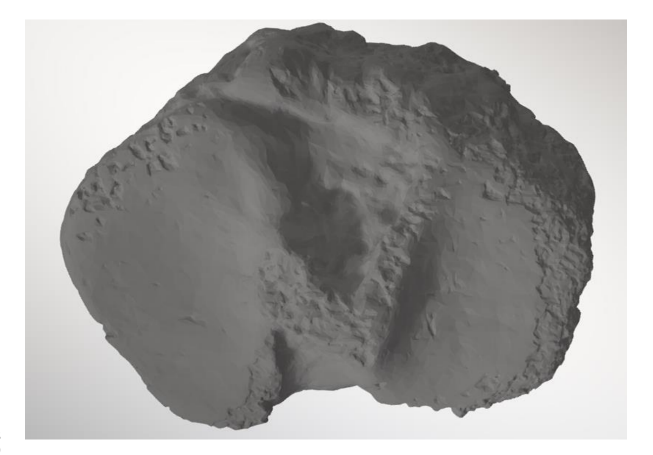


1393 Figure 14: Valgus angles of the knee in, left to right: KNM WT 15000, AL-288-1 and StW

1394 573



- 1402 Figure 15: Proximal tibial surface of Kanapoi KNM-KP 29285A
- 1403 (downloaded from open source: <u>www.africanfossils.org</u>,
- 1404 XYZ dimensions 68.00; 103.30; 60.66 mm)



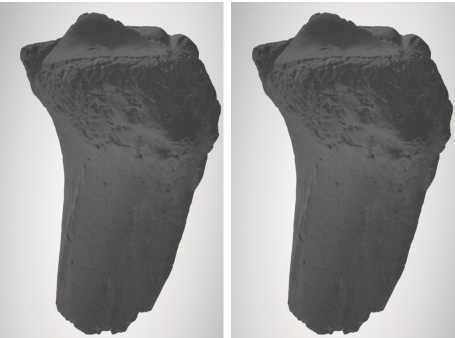
1415 Figure 16: Proximal tibial surface of (Top) StW 573 and (Bottom) KNM WT 15000





- 1424 Figure 17 (Top) Frontal, (Bottom left) Lateral, and (Bottom) Medial perspectives of
- 1425 Kanapoi KNM-KP 29285A (downloaded from open source: <u>www.africanfossils.org</u>, XYZ
- 1426 dimensions 68.00; 103.30; 60.66 mm)





1436 Figure 18 (Top) Frontal, (Middle) Lateral and (Bottom) Medial perspectives of proximal

tibial condyles of StW 573

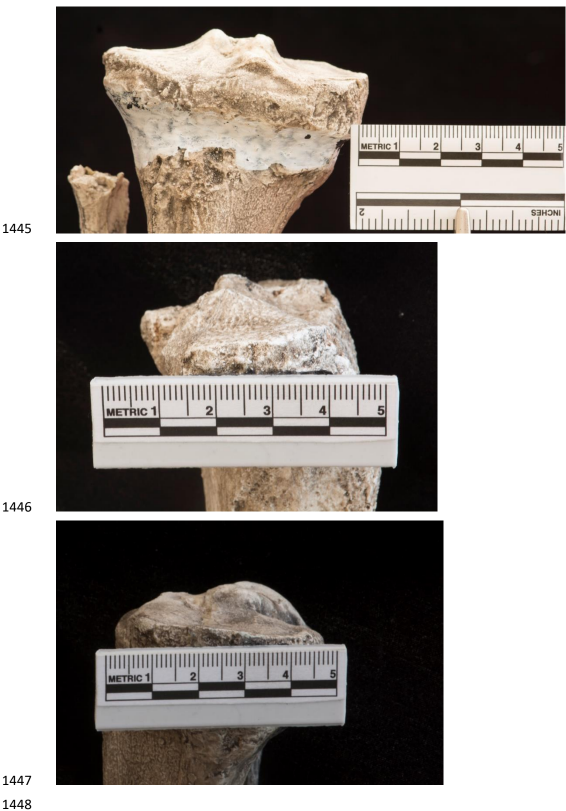




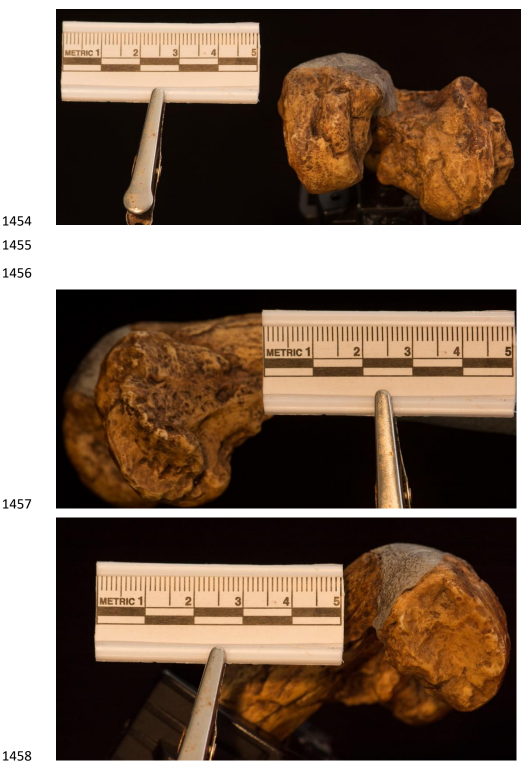


Figure 19 (Top) Frontal, (Middle) Lateral and (Bottom) Medial perspectives of proximal

tibial condyles of KNM WT 15000



- 1450 Figure 20 (Top) Axial, (Middle) Lateral and (Bottom) Medial perspectives of distal femoral
- 1451 condyles of AL-288-1



- 1461 Figure 21 Radius of curvature of the distal tibial condyle/plafond
- 1462 Top: StW 573 Bottom: KNM WT 15000





- 1469 Figure 22 Kanapoi distal right tibia KNMKP 29285 from lateral side, ((downloaded from
- 1470 open source: <u>www.africanfossils.org</u>,
- 1471 Dimensions: x=40.16; y=97.82; z=40.50 mm.)
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- 1473
- 1474 1475
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- 1477 Figure 23 Long bones of the upper and lower limbs of StW 573



- 1482 Figure 24 Archival image of an indigenous arboreal forager climbing a thin vine using flexed
- 1483 elbow postures and hallucal grasp (courtesy of Kirk Endicott)



- Figure 25: An indigenous Batek arboreal forager demonstrating his hallucal grasp for
- climbing a small vine (video frame, courtesy of Vivek Venkataraman)

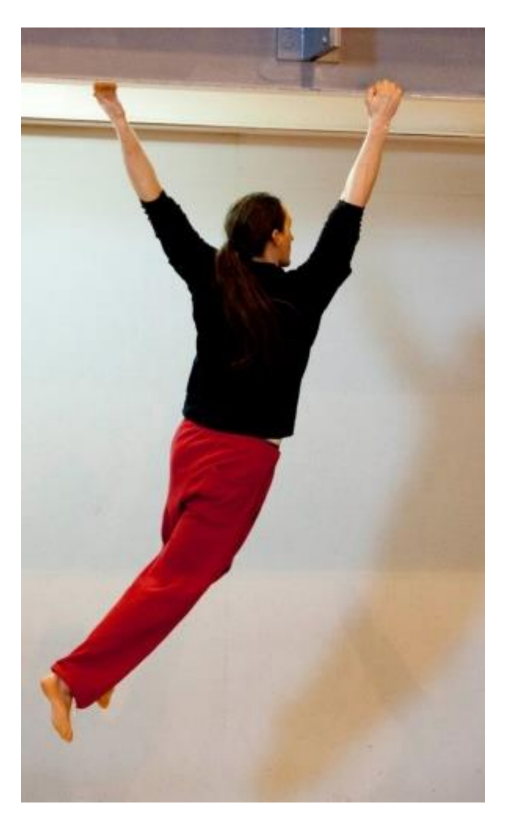


- 1498 Figure 26: Frames from the official video (<u>'Drawin' on Darwin' Live drawing created at the</u>
- 1499 <u>Meeting of the Primate Society of Great Britain | Saranjit Birdi | Axisweb)</u> of: Top, the artist
- 1500 Saranjit Birdi drawing with his foot Bottom: closeup to show precision of hallucal grasp
- 1501 (courtesy of Emily Saunders and the artist);
- 1502



1503

- 1506 Figure 27: Video frame of a parkour athlete brachiating on an I-beam, demonstrating the
- 1507 plasticity of human finger capabilities (see Halsey et al. 2017, permission of the subject and
- 1508 courtesy Susannah K.S. Thorpe)
- 1509
- 1510



- 1512 Figure 28: Handbones of StW 573



- 1518 Figure 29: Apical ridge on the trapezium of: left *Gorilla gorilla beringei*, right StW 573
- 1519 (originals)

