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Musculoskeletal modeling of an ostrich (*Struthio camelus*) pelvic limb : Influence of limb orientation on muscular capacity during locomotion

We developed a three-dimensional, biomechanical computer model of the 36 major pelvic limb muscle groups in an ostrich (*Struthio camelus*) to investigate muscle function in this, the largest of extant birds and model organism for many studies of locomotor mechanics, body size, anatomy and evolution. Combined with experimental data, we use this model to test two main hypotheses. We first query whether ostriches use limb orientations (joint angles) that optimize the moment-generating capacities of their muscles during walking or running. Next, we test whether ostriches use limb orientations at mid-stance that keep their extensor muscles near maximal, and flexor muscles near minimal, moment arms. Our two hypotheses relate to the control priorities that a large bipedal animal might evolve under biomechanical constraints to achieve more effective static weight support. We find that ostriches do not use limb orientations to optimize the moment-generating capacities or moment arms of their muscles. We infer that dynamic properties of muscles or tendons might be better candidates for locomotor optimization. Regardless, general principles explaining why species choose particular joint orientations during locomotion are lacking, raising the question of whether such general principles exist or if clades evolve different patterns (e.g. weighting of muscle force-length or force-velocity properties in selecting postures). This leaves theoretical studies of muscle moment arms estimated for extinct animals at an impasse until studies of extant taxa answer these questions. Finally, we compare our model's results against those of two prior studies of ostrich limb muscle moment arms, finding general agreement for many muscles. Some flexor and extensor muscles exhibit self-stabilization patterns (posture-dependent switches between) flexor/extensor action) that ostriches may use to coordinate their locomotion. However, some conspicuous areas of disagreement in our results illustrate some cautionary principles Importantly tendon travel empirical measurements of muscle moment arms

must be carefully designed to preserve 3D muscle geometry lest their accuracy suffer relative to that of anatomically realistic models. The dearth of accurate experimental measurements of 3D moment arms of muscles in birds leaves uncertainty regarding the relative accuracy of different modelling or experimental datasets such as in ostriches. Our model, however, provides a comprehensive set of 3D estimates of muscle actions in ostriches for the first time, emphasizing that avian limb mechanics are highly threedimensional and complex, and how no muscles act purely in the sagittal plane. A comparative synthesis of experiments and models such as ours could provide powerful synthesis into how anatomy, mechanics and control interact during locomotion and how these interactions evolve. Such a framework could remove obstacles impeding the analysis of muscle function in extinct taxa.

Musculoskeletal modeling of an ostrich (*Struthio camelus*) pelvic limb: Influence of limb orientation on muscular capacity during locomotion

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32 Introduction

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34 As the largest living avian bipeds, ostriches (Struthio camelus Linnaeus 1758) are important for understanding how body mass influences locomotor mechanics in birds. In 35 36 addition, ostriches are among the fastest of living terrestrial animals, and are the fastest living (perhaps even the fastest ever) bipedal runners. These birds can reach maximum speeds >15 37 38 ms⁻¹ (Alexander et al., 1979); similar to another biped that is coincidentally of similar size: 39 red kangaroos (Macropus rufus) (Bennett & Taylor, 1995). Examination of their locomotor 40 dynamics may reveal some of the complex factors that determine maximum running speed in 41 land animals and guide the development of fast running machines. Ostriches are also of 42 similar body size to humans, which other than birds are the only obligate striding bipeds 43 today, making comparisons of bipedal locomotor function in these two species possible (e.g., 44 Gatesy and Biewener, 1991; Rubenson et al., 2011). Additionally, as the largest extant birds, 45 ostriches are important "endpoints" for studies of body size effects on locomotion (e.g., Maloiy et al., 1979; Gatesy et al., 2009; Brassey et al., 2013a,b; Kilbourne, 2013). 46 Furthermore, ostriches are members of the ratite bird clade, whose evolution from basal flying birds into large cursorial flightless animals has been of longstanding scientific interest. However, the evolutionary patterns and processes that produced the diversity of living ratites and their unusual locomotor mechanisms remain uncertain (Baker et al., 2014 and references therein). In turn, ratite birds including ostriches occupy relatively basal positions in extant avian phylogeny (e.g., Clarke and Cracraft, 2001; Baker et al., 2014). Despite their remarkable cursorial specializations and evolutionarily increased body size, ostriches can offer clues to the structure and function of earliest crown clade birds, and thereby about the evolution of avian locomotion from their theropod dinosaur forebears (e.g., Gatesy, 1990). Ostriches have also often been employed as analogues for dinosaur locomotion (e.g., Russell, 1972; Paul, 1998) despite some major anatomical differences (Gatesy, 1990, 1995; Hutchinson and Gatesy, 2000; Hutchinson, 2002; Gatesy et al., 2009; Hutchinson & Allen, 2009), so ostriches and extinct dinosaurs may be reciprocally informative.

A general problem facing those interested in examining the above questions in detail 61 is that ostrich locomotion, including pelvic limb structure-function relationships during 62 movement, remains incompletely understood. Numerous studies have empirically 63 investigated the locomotor kinematics and kinetics of ostriches (e.g., Alexander et al., 1979; 64 Gatesy and Biewener, 1991; Abourachid, 2001; Abourachid and Renous, 2000; Rubenson et 65 al., 2004,2007,2011, Smith et al., 2006,2007,2010,2013; Jindrich et al., 2007; Schaller et al., 2009,2011) and much focus has been given to the whole-body mechanics and energetics of 66 67 ostriches (e.g., Fedak and Seeherman, 1979, Fedak et al., 1982; Rubenson et al., 2004, 68 Watson et al., 2011). However, detailed understanding of the interactions between the various 69 components of the musculoskeletal system and the environment during these movements 70 remain poorly understood. Due to the complex, non-linear dynamics of the musculoskeletal 71 system, correlating whole-body level measures of locomotion to specific muscle function-72 structure relationships cannot vet be adequately performed.

73 Furthermore, ostrich myology was not carefully described until recently 74 (Weissengruber et al., 2003; Gangl et al., 2004; Zinoviev, 2006; also knee joint functional 75 morphology by Fuss, 1996; foot/ankle function by Schaller et al., 2009,2011). Previous 76 myological studies were marred by errors in interpreting ostrich anatomy and by confusing application of mammalian anatomy to ostriches (e.g., Haughton, 1864; Macalister, 1864) or 77 78 provided only a superficial treatment of proximal pelvic limb anatomy (e.g., Mellett, 1994). 79 Even Gadow (1880), a classic comparative reference, contains antiquated concepts of 80 homology (Rowe, 1986). These prior studies of ostrich myology augment comparative work on the pelvic limb myology of other ratites (e.g., Haughton, 1867a,b; Gadow, 1880; 81

McGowan, 1979; Vanden Berge, 1982; Patak and Baldwin, 1998; Picasso 2010,2012; Lamas et al., 2014; Regnault et al., 2014). Additionally, data are available on muscle physiology in ostriches (Velotto and Crasto, 2004) and other ratites (e.g., McGowan, 1979; Patak and Baldwin, 1993), although biomechanical data characterizing muscle force-velocity and forcelength relationships for avian pelvic limb muscles are scant (e.g., Nelson et al., 2004).

This body of prior research provides the strong foundation necessary for detailed 87 88 examination of ostrich limb muscle function using anatomically-realistic biomechanical 89 modelling in order to advance understanding of how the largest living bird supports its body 90 weight and moves itself with its pelvic limb muscles. Here, to provide new insight into 91 ostrich locomotor structure-function relationships, we investigate how pelvic limb muscle 92 functions o relate to limb orientation (i.e., posture/pose or joint angles; Gatesy, 1995; Gatesy 93 et al., 2009) in ostriches. To do this, we integrate data from experimentally measured joint 94 kinematics and ground reaction forces with a biomechanical computer model that was 95 constructed by digitizing the bones, muscles, and tendons of an adult ostrich. By replicating 96 ostrich structure and behavior, the musculoskeletal model provides estimates of individual muscle moment arms (Pandy, 1999) and maximum capacity for moment generation during those behaviors: quantities that are otherwise difficult or impossible to accurately measure non-invasively, especially for all thirty-six major pelvic limb muscles simultaneously. Here we use this model to address some fundamental mysteries about locomotion in ostriches as well as birds, non-avian dinosaurs and bipeds.

Larger mammalian species tend to have straighter limbs than smaller species to improve their effective mechanical advantage during movement (Biewener, 1989, 1990). Yet it remains unclear how much of this improvement is achieved by reducing the moment arms of ground reaction forces about the limb joints using less flexed limb joint orientations or by increasing muscle moment arms via increased anatomical leverage (e.g., relatively larger trochanters) or straightened limb orientation (i.e., which shifts muscles further away from joint centres; Hutchinson et al., 2005). Our ostrich musculoskeletal model will facilitate discerning this relationship in birds, for whom it seems a similar pattern to mammals of having improved mechanical advantage in larger species holds (Gatesy and Biewener, 1991; Hutchinson, 2004; Günther et al., 2004; Brassey et al., 2013; Kilbourne et al., 2013).

112 Previous studies (e.g., Hutchinson et al., 2005; and references therein) suggested that 113 limb antigravity muscle moment arms (or moment-generating capacity; Full and Ahn, 1995) 114 may peak in very upright limb orientations, which intimates that smaller animals with more crouched poses (e.g., birds) employ sub-optimal joint angles for supporting their body weight 115 (Biewener, 1989; Günther et al., 2004), presumably as a tradeoff to provide other benefits 116 117 such as increased maneuverability (e.g., Daley and Usherwood, 2010). As the largest living birds, ostriches stand and move with straighter limbs than smaller birds (Gatesy and 118 119 Biewener, 1991). However ostriches are bipeds that still habitually support themselves with 120 markedly flexed hip and knee joints, which make them a useful case study of this mechanical 121 relationship between posture and antigravity muscle capacity. Understanding this relationship 122 impacts the broader question of why animals choose certain postures—do they select postures 123 that favour larger moment arms for economical force production (e.g., Fujiwara, 2009; 124 Fujiwara et al., 2011; Fujiwara and Hutchinson, 2012) or other factors such as muscle force-125 length properties (e.g., McClearn, 1985; Lieber & Boakes, 1988a,b; Lieber & Brown, 1992; 126 Lieber & Shoemaker, 1992; Lieber, 1997)? 127 The moment arms of ostrich pelvic limb muscles have been studied before with two

different methodologies: two-dimensional (2D) experimental ("tendon travel") measurements

of defleshed limbs (Smith et al., 2007) and 3D musculoskeletal computer models constructed
 from literature data, dissections and scanned skeletons (Bates and Schachner, 2012). The

131 question remains open, how accurate are these experimental measurements and models, and

132 how consistently can different researchers construct such models given the inherent 133 subjectivity involved? This methodological question impacts many of the questions above; a 134 weak model impairs the ability to test hypotheses. Here, we use our 3D musculoskeletal 135 model and previously collected experimental data to address three main questions related to 136 the issues described above: (1) Do ostriches adopt limb orientations during walking or 137 running that optimize their capacity to generate maximal moments about the pelvic limb 138 joints? (2) Are the moment arms of limb muscles maximized (for antigravity/extensor 139 muscles; or minimized in the case of antagonistic flexor muscles) at mid-stance of 140 locomotion (optimizing weight support), or at highly extended limb orientations, as prior 141 studies of dinosaurs inferred (e.g., Hutchinson et al., 2005)? Finally, (3) how accurate or 142 repeatable are estimates of limb muscle moment arms in ostriches using different methods? 143 We also integrate our results with previous studies of ostriches and other large birds 144 (cited above) to infer how the pelvic limb muscles function in locomotion. This provides a

(cited above) to infer how the pelvic limb muscles function in locomotion. This provides a three-dimensional perspective on avian musculoskeletal function, a necessary shift away from past planar (2D) simplifications of this system (e.g., Alexander et al., 1979; Abourachid, 2001; Hutchinson, 2004; Gatesy et al., 2009) because recent studies have shown complex 3D limb dynamics during avian movement (Gatesy, 1994, Hutchinson and Gatesy, 2000; Rubenson et al., 2007,2011; Goetz et al., 2008; Abourachid et al., 2011; Andraka et al., 2013; Kambic et al., 2014). Finally, we synthesize our results with similar data from other bipeds, including humans and *Tyrannosaurus rex*, to infer how limb muscle moment-generating capacity more generally relates to limb orientation and body size.

Materials and Methods

Animals

Experimental biomechanics data were collected from three female adult ostriches 158 159 (70.0, 78.7, 75.9 kg body mass), from which a single representative animal's data was used as 160 model input (78.7 kg). Birds were housed in a large outdoor paddock (5000 m^2) and provided 161 with unlimited access to food and water. All experiments were performed in accordance with 162 the Animal Ethics Committee of the University of Western Australia. Architectural and 163 geometric data for the muscle-tendon units, and skeletal dimensions used in the computer 164 model, were measured in a third female ostrich (65.3 kg), which had no musculoskeletal 165 pathologies and was culled from a commercial ostrich herd (The Ostrich Meat Company, Merced, California). We subsequently verified these anatomical data by qualitative 166 167 comparisons with three other adult specimens of qualitatively similar size, as well as 168 literature descriptions (Gangl et al., 2004; Zinoviev, 2006).

169170 *Kinematic measurements*

171 The experiments and 3D kinematic analyses are described in detail elsewhere 172 (Rubenson et al., 2004,2007,2011). Briefly, the birds were trained to walk and run across a 173 50m long fenced runway surfaced with high-density rubber matting (10 mm thickness). An 174 eleven-parameter direct linear transformation (DLT) was used to construct a 3D image volume from two high-speed cameras (200 Hz) positioned at 45° angles to the runway (Peak 175 176 Motus; Peak Performance, Centenial, CO). The DLT was calculated using a custom-built 48-177 point moveable calibration frame. The video footage from the two cameras were gen-locked and synchronized manually using a hand held switch that caused a barcode in each video 178 179 field to turn white. All data collection was performed after sunset under artificial lighting. A wide range of speeds was recorded in prior studies (Rubenson et al., 2004,2007,2011), but we 180 only used representative kinematic data for a single walking at 1.22 ms⁻¹ and slow running 181

(at 3.46 ms⁻¹). These two trials were within 1 S.D. of the mean kinematic data for trials from
Rubenson et al. (2007).

184 The 3D position and orientation of the ostrich limb segments were determined by 185 videotaping clusters of non-linear, retro-reflective markers placed on the pelvis, femur, tibiotarsus, and tarsometatarsus and a single marker placed on the end of the third phalanx. 186 These markers were used to define segment rigid-body Technical Coordinate Systems (TCS; 187 188 Cappozzo et al., 1995). Prior to walking and running trials, static calibration trials were 189 performed in order to identify several key anatomical landmarks necessary to construct 190 segment Anatomical Coordinate Systems (ACS; see Rubenson et al. (2007,2011) for 191 details). Anatomical landmarks were identified using a 6-marker pointer device and expressed 192 in the segment TCSs. The static calibration trials thus allowed for the reconstruction of each 193 segment ACS across walking/running strides using the motion data of the segment marker 194 clusters alone.

Three-dimensional marker trajectory data were filtered using a fourth order zero-lag Butterworth low-pass filter (4-12 Hz) and compiled in c3d format (Motion Lab Systems, USA). Kinematics were computed using BodyBuilder modelling software (Oxford Metrics; Oxford, UK). Joint angles were calculated by determining the Euler angles associated with the transformation between the ACSs of the proximal and distal segments of a joint (Grood and Suntay, 1983).Segment ACSs in both the model (below) and in the experimental animals were constructed using the same landmarks (anatomical landmarks and numerically derived axes) and same ordered set of rotations between proximal and distal ACSs. Thus, 3D joint motion was defined equivalently in the musculoskeletal model and in the experimental animals and we were able to use experimentally derived joint angles as input into our musculoskeletal model to estimate muscular mechanics *in vivo* during walking and running.

3D coordinate systems for anatomical dissection

We used a Polaris optical tracking system (Northern Digital Inc., Waterloo, Ontario) to record the positions of anatomic landmarks and relevant joint kinematics during dissection sessions. This tracking system is accurate to within 1.5 mm with the 1.5 m³ measurement volume used in this study (Traxtal Inc., Toronto, Ontario), and in order to ensure this accuracy we performed appropriate calibrations before collecting our data.

214 We first skinned the right pelvic limb of the ostrich specimen. Before dissection of 215 the muscles, we attached LED-emitting reference frames (AdapTrax trackers, Traxtal Inc., 216 Toronto, Ontario) to each bone segment using orthopaedic bone screws. Each reference frame 217 contained a cluster of LEDs that allowed the tracking system to record the 3D position and orientation of each segment (establishing the segment TCSs for the dissections, comparable 218 219 to that for the experiments). Figure 1 shows the apparatus we used. We used a digitizing 220 probe (Northern Digital Inc., Waterloo, Ontario) to digitize the 3D coordinates of the 221 musculoskeletal geometry in each session relative to these trackers. Unlike the LED-emitting 222 reference frames, the digitizing probe had a cluster of highly reflective spheres, making it an 223 untethered and mobile tool. When these spheres were visible to the tracking system, the 3D position of the tip of the probe (calibrated in advance) could be recorded with respect to the 224 225 TCS. Three rigid permanent points (marked with a drill as points on the bones) were 226 measured on each segment to provide a local bone coordinate system for all 227 digitizing/dissection sessions. This step allowed the TCS to be removed from the bone and 228 reattached in a different area to facilitate the dissection process while still preserving the

229 overall relationship of digitized points on a given bone between sessions.

230 Building a musculoskeletal model required points to be expressed in the segment 231 ACSs (Figure 2 and Rubenson et al., 2007,2011). The pelvis reference frame was defined as follows: the origin at the midline of the pelvis halfway between the left and right side hip 232 233 joint centres; the unit vector IL SUL (x-axis; positive being anterior); the cross-product of the 234 x-axis and the unit vector SUL SYN (y-axis; positive being cranial), and cross-product of the 235 x-axis and y-axis (z-axis). To locate the hip joint centres, we digitized 10-20 points in and 236 around the acetabulum and femoral head, and then used least-squares optimization to fit a 237 sphere to each of the two resulting point clouds. The centre of this best-fit sphere was the hip 238 joint centre. To establish the reference frames for the other segments, we first estimated the 239 medial-lateral joint rotational axis for the remaining joints by flexing and extending each 240 joint and recording the 3D position and orientation of the distal bone with respect to the proximal one as a series of homogeneous transformation matrices. With these transformation 241 242 matrices, we were able to calculate the average kinematic screw (helical) axes (Bottema and 243 Roth, 1990) that best approximated the flexion-extension axis between those segments.

The femur coordinate system was defined as: the origin at the proximal joint centre; the segment z-axis along the medial-lateral joint rotational axis (positive being lateral); the y-axis as the cross-product of the z-axis and the unit vector between the proximal and distal joint centers; and the x-axis as the cross-product of the y- and x-axes. The tibiotarsus and tarsometatarsus coordinate systems were defined as: the origin at the proximal joint centre; the y-axis as unit vector between the proximal and distal joint centre; the y-axis as unit vector between the proximal and distal joint centers; the segment z-axis as the cross product of the medial-lateral joint rotational axis and the y-axis; and the x-axis as the cross-product of the y- and z-axes. The pes coordinate system was defined as: the origin at the proximal joint centre; the segment x-axis as the unit vector between the proximal joint centre; the origin at the proximal joint centre; the segment x-axis as the unit vector between the proximal joint centre; the segment y-axis as the unit vector between the proximal joint centre; the segment x-axis as the unit vector between the proximal joint centre; the segment; the z-axis as the unit vector between the proximal joint center and the end of the segment; the z-axis as the cross-product of the x- and z-axes. Putting any digitized points into these ACSs required two linear transformations: from the TCS into the local bone coordinate system and subsequently into the ACS. Table 1 provides data on axis positions used in the final model.

Anatomical digitization and musculoskeletal model construction

260 We began by dissecting the specimen proceeding from superficial to deep structures. 261 The positions of muscles and bone geometry or other relevant anatomical features on each limb segment were measured using the digitizing probe. In particular, before removing 262 muscles we digitized the circumferences of muscle origins and insertions as well as the 3D 263 264 paths of the muscles from origin to insertion, using from 1-30 (depending on extent of the 265 structure) x,y,z coordinate points to characterize each structure of interest. In addition, we measured other musculoskeletal features used as references, such as bone surfaces (for later 266 alignment of complete 3D bone images from CT data), condylar contours, and those 267 268 ligaments that influence muscle-tendon unit paths. All 3D points for use in the model were 269 converted into the ACS.

270 After dissection the bones were defleshed and macerated, but with articular cartilages 271 and menisci remaining as intact as possible. The bones were then CT scanned (1.5-3 mm 272 slices, 120-130 kPa, 109-150 mA, on a Picker PQ5000 CT scanner) and the resulting slices 273 digitized in Mimics software (Materialise, Inc; Leuven, Belgium) to produce rendered 3D polygonal meshes. Ultimately ASCII (ASC) format images were used (decimated to <50,000 274 275 polygons each) as bone images in the model. Importantly, the bones were only visual aids 276 and not inherent obstacles to joint or muscle motion, but they are crucial for visualizing 277 musculoskeletal function.

We used these anatomical data and bone images to construct a 3D musculoskeletal computer model of the right pelvic limb using SIMM software (Musculographics, Inc.;

280 Chicago, IL; Delp et al., 1990,1992; Delp and Loan, 1995,2000). The right limb was mirrored as a left limb. The digitized muscle paths were used as a template for developing the muscle 281 282 paths in the final model (Figures 3, 4). Paths were represented using a combination of "via 283 points" (i.e., static points fixed relative to a segment) and "wrapping surfaces" that prevented 284 translation of points outside of a predefined area (see Delp and Loan, 1995, 2000 for details; 285 also Hutchinson et al., 2005, for a similar procedure we used for Tyrannosaurus rex). Table 2 286 presents the muscles modelled, with abbreviations used throughout the paper. Table 3 287 describes all the muscle wrapping surfaces assumed in the model and Figure 5 displays 288 examples.

Inevitably, because the 3D musculoskeletal geometry was complex, we had to cautiously judge where to position wrapping surfaces and what size and shape they should be. Additional ostrich cadaveric material was used along with the literature (Gangl et al., 2004; Zinoviev, 2006) to qualitatively refine the model as we iteratively progressed, checking that paths and attachments were represented reasonably and consistently. We took care to visualize the model in many different 3D joint positions to ensure that muscles did not pass through areas occupied by other soft tissues or especially bones and to eliminate other numerical errors generated by interactions of the muscle-tendon unit paths with wrapping surfaces (e.g., "loops" in muscles caused by contradictory constraints in the model). Importantly, because we intended to compare our model's results with data from Smith et al. (2007) and Bates and Schachner (2012), we kept our model construction blind to the results of these studies, avoiding any comparisons and indeed finishing the major steps in completing our model before these studies were published.

Muscle-tendon unit architecture and physiology

After we dissected, digitized, and removed the muscles, we separated them from their proximal/distal tendons and other connective tissue. We then used digital calipers (± 0.1 mm), an electronic balance (± 0.001 g), and a protractor ($\pm 1^{\circ}$) to measure muscle fascicle lengths (*L*), masses (m_{musc}), and resting pennation angles (θ) for calculating physiological crosssectional area (A_{phys}), taking an average of five randomized measurements for *L* and θ in larger muscles.

Using water displacement (immersing sectioned muscles in graduated cylinders) to calculate muscle belly (sans tendon) density (*d*) from (volume m_{musc} ⁻¹), we obtained a mean value of 1.0645×10^3 (n=10; S.D.=0.0347) kg m⁻³, matching measurements of mammalian muscle (Mendez and Keys, 1960; Brown et al., 2003a). Hence we assumed a conventional value of *d* as 1.06×10^3 kg m⁻³. As commonly practiced, we assumed *L* to be equivalent to optimal fiber length (l_o^m ; Zajac, 1989). We could thus calculate A_{phys} as (e.g., Alexander et al., 1979; Lieber & Boakes, 1988a; Brown et al., 2003b; Hutchinson, 2004):

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 $A_{phys} = m_{musc} \cos \theta \, (L \, d)^{-1} \qquad (\text{Equation 1})$

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- Equation 2 then estimates F_{max} , maximum isometric force capacity:
- $F_{max} = 3.0 \times 10^5 \text{ m}^{-2} A_{phys} \qquad (\text{Equation 2})$

In which the constant is isometric stress under maximal activation (Medler, 2002; Nelson et al., 2004). Note that the musculoskeletal model varies θ with *L* to maintain constant muscle thickness (Zajac, 1989). Muscle maximal contraction velocity was not calculated in the present model, but could be added (see Smith et al., 2006 for example) for more complex simulations, using published fibre type data for ostriches (Velotto and Crasto, 2004) and ostrich or other avian muscle force-velocity data (e.g., Nelson et al., 2004). We also did not 331 (Zajac, 1989) or specific measurements can be added to future simulations. For simplicity, we 332 chose to focus in this initial study on near-isometric muscle action and progress to more 333 complex, dynamic parameters in later work. 334 Muscles were identified as in Table 2 following Gangl et al. (2004) and Zinoviev 335 (2006) (see Appendix 1 for additional details). For simplicity, we combined some muscle 336 heads into single functional muscles in the model (cf. Gangl et al., 2004:table 1) and omitted 337 some tendinous/minute muscles which were grouped with other similar muscles nearby or

338 omitted in the case of M. popliteus (rotates fibula around tibia; Fuss, 1996); these 339 simplifications are outlined in the Supplementary Text. However, some muscles (e.g., M. 340 iliofibularis, M. iliotrochantericus caudalis) were large enough that separation into two heads 341 was deemed important, as some heads might have very different flexor/extensor moment 342 arms than more cranial/caudally-positioned ones.

include measurements of tendon force-length data here, but either dimensionless estimates

Limb muscle biomechanics: Calculations and hypothesis testing

The musculoskeletal model was then imported into OpenSim (opensim.stanford.edu) software in order to take advantage of the programme's established analysis capabilities. OpenSim uses the 'virtual work' method (change of muscle-tendon unit length per unit joint rotation) explained by Delp and Loan (1995, 2000) and Pandy (1999) to compute muscular moment arms over a range of motion. Maximal muscular moments then can be estimated using muscle F_{max} and potentially l_o^m (see above and Zajac, 1989).

To test whether ostrich muscle moment-generating capacity is optimized to match peak loads during walking and running (our Question 1), we compared the results from estimated maximal muscle moments to experimentally-calculated internal and external moments (Rubenson et al., 2011), addressed in the Discussion. First, each muscle's maximal isometric muscle force (F_{max}) was multiplied by the flexor/extensor moment arm calculated by OpenSim, for each pose adopted throughout the representative walking and running gait cycle trials (every 1% of gait cycle) to obtain the relationship between locomotor kinematics 358 and isometric muscle moments. Second, OpenSim was used to calculate individual muscle 359 moments directly, taking into account muscle force-length relationships (set as dimensionless 360 in a Hill model as per Zajac, 1989), in order to provide a more realistic estimate of the variation of maximal moment-generating capacity throughout the same gait cycles. Both approaches were static, ignoring time/history-dependent influences on muscles. The second 362 approach allowed non-isometric muscle action to be represented, but did not incorporate 363 364 force-velocity effects, which would require a more dynamic simulation to resolve. Total 365 extensor and flexor maximal moments were calculated in OpenSim as well as the net 366 (extensor + flexor) maximal moment.

To determine if ostrich limb muscle moment arms peak at extended limb orientations 367 368 or at mid-stance of locomotion (our Question 2), we used the model to calculate the mean 369 moment arm of all extensor or flexor muscles across the full range of motion of each joint in 370 flexion/extension (set at constant values for midstance of running in other degrees of 371 freedom), summed these mean moment arms, and divided that sum by the summed maximal 372 moment arms for each muscle across the same range of motion (as in Hutchinson et al., 373 2005). We then inspected whether our representative mid-stance poses in walking or running 374 matched maximal or minimal averaged moment arms corresponding to those poses.

375 To compare the degree of matching between muscle moment arms in our model and 376 the experimental data of Smith et al. (2007) and Bates and Schachner (2012) (our Question 377 3), we obtained the published experimental and modelling data (K.T. Bates, provided by 378 request), transformed their joint angle definitions to be consistent with our model definitions, 379 and plotted the muscle moment arms vs. each joint angle with our moment arm data,

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380 restricting the other studies' ranges of motion to those presented in the original studies. For

381 the knee and joints distal to it, in this study we focus only on flexor/extensor moment arms for simplicity and because the importance of long-axis and ab/adduction muscle (vs. passive 382

383 tissue) moments at these distal joints is unclear, although our model could be adjusted to

384 calculate those non-sagittal moment arms and moments.

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387 **Results**

388 Here we present our data for addressing our main questions, proceeding in order with 389 maximal muscular moments, maximal/minimal moment arms, and then general moment arm 390 patterns compared with other studies'. The model is downloadable from the repository at 391 https://simtk.org/home/opensim and can be manipulated in open source software OpenSim. 392 Supplementary Movie S1 shows the model animated through the representative running 393 stride. 394

Maximal muscular moments

Our 3D ostrich limb model predicted how the maximal capacity to generate muscle moments should vary with limb orientation during walking and running (Figures 6, 7). Maximal flexor moments increase if force-length properties are ignored (treating all muscles as isometric). This indicates that most muscles in the model are at disadvantageously short fibre lengths during locomotion, with walking having a generally greater capacity for flexor moment generation (especially about the hip) than running. These curves do not change much across the gait cycle. The pattern for extensor moments is more complex. Peak capacity tends to be in late swing phase (reasonably consistent across all joints). Force-length properties here provide an advantage, presumably because the muscles are lengthened. Data during the stance phase do not support the hypothesis, regardless of assumptions about muscle-force length states, that postures used around mid-stance of walking or running optimize the moment-generating capacity of pelvic limb muscles in ostriches: the maximal moments early or late in stance phase, and late in swing phase, are of similar or greater magnitudes. The relatively flattened shapes of most moment curves without force-length properties enforced ("Fmax"; dotted lines in Figures 6,7) indicate that muscle moment arm variation across postures used in vivo during locomotion is a smaller contributor to moment generation than force-length properties ("F-L"; solid lines) in Struthio.

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414 Maximal/minimal muscle moment arms and limb orientation

Do ostriches' limb muscle moment arms peak at very extended limb orientations or at 415 midstance of walking/running (Figure 8)? We find that the mean hip extensor moment arms 416 417 decrease from a peak at full extension as hip joint flexion increases, and the hip flexors 418 behave similarly. However, knee and ankle moment arms each exhibit different patterns. The 419 knee extensor and flexor moment arms tend to peak at moderate knee flexion angles (~60-420 90°), as do the ankle extensors (plantarflexors), but the ankle flexors have a near-plateau for 421 most angles, quickly decreasing with extreme dorsiflexion (>100° ankle angle).

422 When the poses that ostriches use during periods of peak limb loading (near 423 midstance of walking and running; Rubenson et al., 2007) are compared against these 424 patterns (Figure 8), it becomes evident that there is no clear optimization of muscle moment 425 arms for supportive (large extensor or small flexor values) roles during these periods of 426 potential biomechanical constraints. This is in agreement with the maximal moment data 427 from Figures 6 and 7. Hip extensors and flexors as well as ankle extensors are relatively far 428 (~60-85% of maximal mean moment arms) from optimal values at midstance of walking and 429 running. Knee extensor/flexor moment arms are closer to maximal values, especially for

walking. However, the co-contraction of multiarticular hip extensor/knee flexors (e.g. ILFB,
FCLP) against knee extensors would eliminate associated benefits—i.e., the ratio of peak
knee extensor to peak knee flexor moment arms would have not have minimized the net knee
extensor moments required at midstance of either walking or running. At moderate knee
flexion values, both the capacity of muscles to extend and to flex the knee are near-maximal
(Figure 8).

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437 Moment arms: general trends and comparisons with prior studies

Figures 9-11 show our results for hip flexion/extension moment arms of ostrich muscles, with comparable data from Smith et al. (2007) and Bates and Schachner (2012) also plotted if available (abbreviated in this section as S.E.A. and B.A.S. respectively). Here we focus on the major findings. The two AMB muscles (Figure 9) compare reasonably well among all three studies, showing a decrease of hip flexion moment arms at strongly flexed limb poses and in some cases (our AMB1,2 and the AMB of B.A.S.) a switch from flexor to extensor action with flexion (~30-90°). The IC muscles likewise have reasonably comparable results, but only our IC muscle switches action at extreme flexion. Our model agrees well with the data of S.E.A. and especially B.A.S. for the IL muscle, including its decreasing hip extensor moment arm with increasing hip flexion and a switch from hip extensor to flexor action at typical *in vivo* positions (~40-70°). We have similar findings for the ILFB muscle, although no switch to hip flexor moment arms is observed in either of the two parts of this muscle in our model (S.E.A. and B.A.S. represented it as one part) (Figure 9).

Uniarticular muscles acting about the hip joint consistently display flexor action for the IFE, IFI, ISF and OM muscles (Figure 10). We find fair agreement among studies for the IFE (note confusion caused by misidentification of muscles in prior studies-- see Appendix 1; the "IFE - Smith" in Fig. 10 is equivalent to our IFE and ITC), ITC, IFI, ITM and ITCR muscles' general changes of moment arms. Our IFE moment arm values are smaller than for S.E.A. and B.A.S. apparently because of the aforementioned identification issue (the top left panel in Fig. 10 shows our IFE plotted against S.E.A.'s IFE+ITC combined). Notably, the curves for the two parts of ITC in our data and those of B.A.S. are remarkably similar (and 459 consistent with S.E.A.'s experimental data for their "IFE – Smith" as well as "ITC - Smith") 460 despite the subjectivity inherent in partitioning this large muscle into two paths. These 461 moment arms grade from flexor to extensor action with strong flexion (~40-70°). A similar 462 trend is evident for the ITM and ITCR muscles (but note the identification issues outlined in Appendix 1; S.E.A.'s "ITC" is actually the ITM, which their data otherwise lacks, so the top 463 464 right panel in Fig 10 compares their actual ITM ["ITC – Smith"] vs. our ITM). The 465 antagonistic OM and ISF muscles concur less closely between the latter two studies, 466 however, displaying more convex curves tending to indicate hip flexor action in our data, 467 with more concave, flattened arcs favouring hip extensor action in B.A.S. (Figure 10).

468 The "hamstring", caudofemoral and adductor hip muscles uniformly display extensor 469 action, befitting their more caudal paths relative to the hip, but agree less well among studies 470 than the prior muscles (Figure 11). Our data for the FCM, FCLP, CFP and PIFML muscles 471 portray peak moment arms at low hip extension angles ($\sim 0-30^{\circ}$), decreasing with flexion 472 away from these ranges. These trends qualitatively agree with the S.E.A. and B.A.S. data, but 473 moment arm values tend to be substantially smaller in those data, especially for the FCLP and 474 FCM muscles. Our PIFML data show less variation with joint angle than the S.E.A. and B.A.S. data because we had to constrain this muscle's path in 3D to avoid it cutting through 475 476 bones or other obstacles in some poses. Note also how the S.E.A. results in general show 477 strong changes with joint angles, whereas the more constrained muscle geometry of our model and B.A.S.'s results in more modest changes (Figure 11). 478

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479 Long-axis rotation (LAR; in Figures 12,13) moment arms for hip muscles only allow 480 comparisons between our data and those of B.A.S. Furthermore, as B.A.S. plotted these moment arms against hip flexion/extension joint angle, we show them that way here but also 481 482 plot them against hip LAR joint angle in the Supporting Information (Supplementary Figures 483 S1,S2); however we do not discuss the latter results here. For the AMB1,2 muscles we find consistently weak, near-zero LAR action (lateral/external rotation), whereas B.A.S. showed a steeply increasing hip medial/internal LAR moment arm as the hip is extended (Figure 12). In contrast, our IC muscle data agree well with B.A.S.'s in having a shallow increase of the medial/internal LAR moment arm with hip flexion. The two heads of the IL muscle show opposite trends in our results vs. B.A.S.'s (in our data, becoming less effective at lateral/external rotation as the hip is flexed). Our results for the two parts of the ILFB muscle are very different from B.A.S.'s in trending toward stronger medial/internal rotation function, whereas B.A.S.'s favour lateral/external rotation. The results for the OM muscle have a more intermediate quality of matching between studies but still indicate a lateral/external rotation action for this large muscle. Contrastingly, our ISF data and those of B.A.S. match fairly closely, with consistent lateral/external rotator action. The FCM and FCLP muscles have among the largest LAR moment arms for all muscles (~0.08m; also observed for our ILp muscle) in our data, but both muscles reduce their lateral rotator action with increasing hip extension. In B.A.S.'s data a weaker, opposite (medial/internal rotator) trend was found for these muscles (Figure 12).

The uniarticular hip muscles' LAR moment arms tend to switch more often from medial to lateral rotation or vice versa (Figure 13). The IFI, however, remains mainly as a weak medial rotator except at extreme hip flexion (>60°). B.A.S.'s data favoured a stronger medial/internal rotation moment arms for the IFI. Our IFE muscle's data indicate a switch from lateral rotation into medial rotator action. Our results for the two-part ITC muscle concur qualitatively with B.A.S.', consistently having a strong medial rotator action but smaller at more extended joint angles. Finally, as in B.A.S.'s data, but featuring smaller moment arms, our data show that the CFP and PIFML muscles have consistent lateral rotator moment arms peak at hip angles of 30-60°, then decrease; a pattern qualitatively matched by B.A.S.'s data. (Figure 13).

511 Abduction and abduction moment arms for the hip muscles show strong postural dependency like the LAR moment arms do (Figures 14-15). Again, as for the LAR data 512 513 above, we supply these data plotted against abduction/adduction hip joint angle in the 514 Supporting Information (Supplementary Figures S3, S4), but we do not discuss those results 515 here. The PIFML muscle has a discontinuity in its hip abductor moment arm (Fig. S4) in our 516 model at extreme hip abduction angles (>-40°) but this is well outside normal in vivo 517 abduction angles used (<25°; Rubenson et al., 2007). The two AMB muscles in our model 518 have peak adductor moment arms at different flexion angles ($\sim 30^{\circ}$ and 80°), then decrease. 519 B.A.S.'s data, in contrast, showed their AMB muscle to act as an abductor. Our IC muscle 520 has a similar adductor moment arm curve as our AMB2, and a similar divergence from 521 B.A.S.'s results. Our IL muscle parts agree reasonably well with B.A.S.'s, showing them to 522 act as adductors. Both our ILFB muscle parts have little variation in their hip abductor 523 actions; B.A.S.'s representation changed steeply and became an adductor with extreme hip 524 flexion. The OM muscle, which runs very close to the plane of the acetabulum, is an adductor 525 at extended joint angles and an abductor at flexed angles in our model, whereas it remained 526 an adductor in B.A.S.'s data. While the ISF muscle is mainly an abductor in our model, it was 527 exclusively an adductor in the B.A.S. model. The FCL and FCM muscles compare only qualitatively between our data and B.A.S.'s, remaining as hip abductors. It is noteworthy that 528

529 throughout the full ranges of hip motion we examined, most muscles would act as hip 530 abductors; the dorsal AMB2 and IC muscles are the only consistently strong hip adductors 531 (Figure 14).

532 Uniarticular "deep dorsal" and antagonistic muscles show similar trends as the above 533 muscles for adduction/abduction capacities (Figure 15). The IFI has weak adductor action, vs. abductor in B.A.S.'s data, whereas our data and B.A.S.'s agree well on the hip abductor 534 535 moment arm of the IFE. Our representations of the ITCa/p muscle parts switch from 536 abduction to adduction function as hip flexion surpasses 45-60°; B.A.S.'s did not. 537 Postacetabular muscles such as the CFP and PIFML in our model are almost exclusively hip 538 abductors, but the caudofemoral muscles of B.A.S. are weak adductors and the PIFML 539 equivalents in that study switched to that same action with flexion. Again, our results are generally opposite B.A.S.'s in the case of the ITM and ITCR muscles, which convert from 540 541 abductor to adductor action at 10-40° hip angles in our model and did not change much in B.A.S.'s (Figure 15). 542 543

We only focused on flexion/extension moment arms for more distal joints, starting with the knee (Figures 16,17). Good agreement between knee extensor moment arms for the AMB2 (dorsal) muscle is evident with the two other studies. The AMB1 (ventral) component only has data from our model (mainly a weak knee flexor), as does the IC (very weak knee flexor/extensor at flexed/extended angles). We estimate a larger knee extensor moment arm for the FMTL muscle but this is because of misidentification of part of that muscle in the S.E.A. and B.A.S. data (only a distal head was included in this muscle; see Appendix 1). Similar differences of anatomical representation are likely explanations for the deviation between our result (weak knee flexor) and S.E.A.'s (strong knee extensor) for the FMTM (see Discussion). Our model presents slightly different moment arms for its two IL muscle heads, peaking in extensor values at 30-90° flexion, whereas B.A.S. had identical moment arms increasing throughout extension.

545 546 547 548 549 550 551 552 553 554 We estimate the knee flexor moment arms as identical (peaking at 90-120° flexion) 555 for the two parts of the ILFB muscle in our model, which match S.E.A.'s data well, whereas 556 557 B.A.S. had moment arms switching from extensor to flexor at 70° of knee flexion, peaking at quite extended knee poses (Figure 16) (see Discussion). Other "hamstring" muscles (no 558 559 comparable data for S.E.A. or B.A.S.) include the FCM and FCLP, which shift steeply from 560 knee extensor to flexor moment arms at high flexion angles and then peak near 90° in its flexor moment arm value (Figure 16). The FL muscle shows an almost mirror image pattern, 561 acting as a knee extensor. Muscles running past the ankle joint (Figure 17), such as the FPD3 562 563 and FPD4 groups, have a similar pattern to the FCM and FCLP at the knee, but the TCf 564 muscle has almost no knee moment arm; consistently acting as a very weak extensor. Finally, 565 parts of the gastrocnemius muscle group (e.g. GIM) reach peak knee flexor moment arms of about 0.07m at intermediate knee flexion angles (60-90°). The data for S.E.A. and B.A.S. and 566 567 for our GL muscle remain(ed) near smaller knee flexor values, with less postural variation 568 (Figure 17).

569 Ankle musculature displays fairly congruent patterns in our model and S.E.A. and 570 B.A.S.'s data (Figures 18,19). The TCf and TCt heads generally have an ankle extensor 571 action, like the EDL muscle group does, albeit with some switches to extensor action with 572 extreme (dorsi)flexion in the B.A.S. dataset (and our TCf). Surprisingly, ankle extensors 573 reveal more variation: our FDL's ankle extensor moment arm is almost twice as large of that 574 in the S.E.A. and B.A.S. data, showing little change with ankle posture, whereas the B.A.S. dataset exhibited a decreased moment arm with flexion. Our other digital flexor muscles (FPD3, FPD4) and those of S.E.A. display roughly similar values but opposite trends, 577 increasing with ankle flexion in our model. Our FL muscle's extensor moment arm is smaller 578 than those of S.E.A. and B.A.S. The model of B.A.S. had a M. fibularis brevis (FB) muscle

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(Figure 18), which is reduced to a ligament in *Struthio* and thus not included in our model; no
studies have data for the ligamentous M. plantaris (Zinoviev, 2006). The extensor moment
arms for our gastrocnemius muscles are all identical and fairly constant with ankle flexion,
whereas the curves for the data of S.E.A. and B.A.S. increased steadily and tended to be
larger (Figure 19).

584 Digital flexor muscle moment arms all stay fairly constant (slight increase with 585 extension of the MTP joint) in our model whereas they showed a stronger decrease in 586 S.E.A.'s experiment (Figure 20). Our EDL muscle has stronger moment arms than in 587 S.E.A.'s data but a similarly shallow curve. Finally, our FL muscle exhibits digital flexor 588 moment arms similar to those of the other digital flexors.

Discussion

The results of our combined experimental and theoretical approach show first that, while ostrich limb muscles are capable of generating large flexor and extensor moments about their limb joints during locomotion (Figures 6,7), they do not seem to match maximal muscle moment-generating capacity with instants of peak loading in walking or slow running. Second, the moment arms of ostrich flexor/extensor muscles often change greatly with limb orientation, but they are not consistently matched to minimize the former and maximize the latter during key periods of weight support in locomotion (Figure 8). Third, there is mostly reasonable consistency in three different studies of ostrich muscle moment arms (Figures 9-20), indicating at least fair repeatability with distinct methods, but still some striking disagreements, especially in the little-explored area of non-flexor/extensor muscle mechanics. We explore these topics in more detail below and then consider broader issues related to our findings.

Maximal muscle moments and kinematics

607 Our Question 1 asked, "Do ostriches adopt limb orientations during walking or 608 running that optimize their capacity to generate maximal moments about the pelvic limb 609 joints?". We find no convincing evidence of such optimization -- maximal capacities to 610 produce joint moments often peak either early in stance phase or during swing phase (Figures 611 6.7). In both cases, net joint moments obtained from inverse dynamics analysis are low (Rubenson et al., 2011). Peak flexor moments (requiring extensor/antigravity muscle activity) 612 613 occur at or near midstance in running ostriches (Rubenson et al., 2011:Figure 7), reaching magnitudes >-2 Nm kg⁻¹ (i.e. >-150 Nm for our subject). Note that these magnitudes are far 614 615 below the capacities of hip, knee and ankle muscles (>400 Nm; Figures 6, 7) but approach 616 those of the digital flexors (~150-200 Nm; Figure 7). On this basis, we infer that either 617 passive tissues (including muscle passive force-length properties) play an important role in 618 balancing moments about the tarsometatarsophalangeal joint in running ostriches (cf. 619 Haughton, 1864; Schaller et al., 2009,2011), especially at faster speeds, or that muscle 620 moment-generating capacity is near its limits for this joint in particular, even at slower 621 speeds. Nevertheless, more proximal limb muscles seem further from their moment-622 generating limits.

In his classic biomechanical analysis of ostrich anatomy, Haughton (1864) assumed that "the greatest possible amount of muscular force shall be expended in straightening or unbending the legs", and thus that early and late stance respectively placed the greatest demands on these forces. Available data no longer support this notion, but there is no question that ostriches have muscle masses able to produce greater moments (and work) in 628 extension than in flexion, as Haughton explained, but by a factor of about three times for the 629 hip and knee rather than ten (*vide* Smith et al., 2006,2007).

630 There are several potential explanations for our observations that lead us to a negative 631 answer to our study's first question. First, we have only examined walking and slow running. Near maximal speed, moment capacity and requirements around midstance might be more 632 closely matched (e.g. Hutchinson, 2004), as forces surely increase. At a duty factor of 0.42, 633 634 Rubenson et al. (2011) obtained peak vertical ground reaction forces of 1500-2000 N or 635 about 2.17-2.89 times body weight (BW), whereas Alexander et al. (1979) estimated 2.7 BW peak forces for an ostrich at near top speed (duty factor 0.29). The latter study used an 636 637 equation that probably underestimates peak forces for ostriches, as Rubenson et al.'s (2011) 638 data show (peak forces are 16-55% greater than predicted from duty factor). Second, our 639 present model is still static, not considering force-velocity or other dynamic interactions that 640 would alter moment-generating capacities. It is possible that these parameters, or highly 641 complex interactions (e.g., muscle moment arms and "power amplification"), could be more influential than the isometric and force-length properties that our model considers. Third, entirely different factors could determine locomotor and postural optimization, such as energetic costs or stability/manoeuvrability (e.g., Daley and Usherwood, 2010).

Comparison of our results with other studies of the relationship between limb orientation and muscle mechanics reveal a fourth potential explanation, that the optimization of anatomy, posture, physiology and other factors in locomotor dynamics could be highly species-, task-, limb-, joint- or muscle-specific. Lieber and colleagues (Lieber & Boakes, 1988a,b; Mai & Lieber, 1990; Lieber & Brown, 1992; Lieber & Shoemaker, 1992) conducted an elegant series of studies that constitute a model system for addressing this issue. They elucidated that maximal moment production by the semitendinosus muscle in frog hindlimbs showed a strong dependence on muscle isometric force capacity and moment arms. Some of these studies found less dependence of moment production on joint angle-dependent moment arm values (e.g., Lieber & Boakes, 1988a,b), but this dependency varied for the hip and knee joints (Mai & Lieber, 1990; Lieber and Shoemaker, 1992)—and might be expected to vary for other muscles, too. Indeed, the moment arm did not vary much with knee joint angle for the semitendinosus (e.g., 0.37-0.44 cm about knee, across 10-160° range of flexion/extension; 658 Lieber & Boakes, 1988a: Figure 6A) so this muscle could not contribute much variation to 659 muscle moment production. One might predict more dependency of maximal muscle moment production on moment arms for muscles that have more variable moment arms, but this has 660 not been conclusively determined. Lieber & Brown (1992) found that there was no simple 661 relationship between muscle fibre length and moment arm in seven frog hindlimb muscles, 662 663 with differences evident between muscles acting about the hip and knee, suggesting diverse 664 adaptations to moment production demands.

The aforementioned studies' reviews of numerous others in humans, cats and other 665 666 species likewise note some variability and uncertainty in what factors determine maximal 667 moment capacity in limbs, so consensus has been elusive. Furthermore, Brown et al.'s (2003b) modelling/experimental study of horse forelimbs found that while the moment-668 669 producing capacities of flexor muscles were determined mostly by muscle properties (e.g., 670 F_{max} or muscle force-length), muscle moment arms could have greater effects on moments 671 than those properties for some extensor muscles. Young et al. (1993) also found variability 672 from 50-100% of resting fascicle length in the amount of length change that cat ankle muscles used throughout their range of motion. Thus strict "sarcomere equivalency" 673 674 (constant usage of maximal muscle range of motion; e.g. discussions in McClearn, 1985; 675 Lieber, 1997; Bates & Schachner, 2012) is not expected, but an approximate "tuning" of moment arms to muscle fiber lengths (and joint ranges of motion) is expected at least in some 676 cases. Lieber and Shoemaker (1992) explained how a greater muscle fibre length to moment 677

678 arm ratio would cause muscle force-length properties to become less influential on muscle 679 moment production. Therefore, in the long-fibred proximal muscles of ostriches and other 680 birds that run closer to the hip and knee joints and thus have smaller moment arms (e.g., 681 Smith et al., 2006,2007), individual muscles' moment arms might be quite influential. This speculation has yet to be conclusively tested, let alone integrated into studies of whole limbs 682 683 and locomotor dynamics. 684

Regardless, recent studies of the hindlimbs of mice (Lieber, 1997), rats (Johnson et 685 al., 2008), and chimpanzees (O'Neill et al., 2013), as well as horse forelimbs (Brown et al., 2003a,b) favour some optimization of locomotor tasks and muscle moment arms, as do broad 686 687 comparative studies of elbow muscles by Fujiwara and colleagues (Fujiwara, 2009; Fujiwara 688 et al., 2011; Fujiwara and Hutchinson, 2012) and a comparative study of mammalian 689 carnivores (McClearn, 1985). So far, however, general principles that extend across lineages, 690 behaviours or anatomies remain elusive. We consider this ambiguity's effect on inferences 691 about extinct taxa further below.

Muscle moment arm-joint angle dependencies

Our Question 2 asked, Are the moment arms of limb muscles maximized (for antigravity/extensor muscles; or minimized in the case of antagonistic flexor muscles) at midstance of locomotion (optimizing weight support), or at highly extended limb orientations, as prior studies of dinosaurs inferred (e.g., Hutchinson et al., 2005)?

The peak extensor muscle moment arms that ostrich pelvic limb antigravity muscles have about the hip joint lie close to a completely columnar (i.e., vertical or 0°) hip angle, approximately 5° (Figure 8), similar to prior results for Tyrannosaurus rex (Hutchinson et al., 2005). Ostriches, however, do not stand or normally move with such extended hip joints (Rubenson et al., 2007). We suspect this difference is because of their two orders of magnitude smaller body size (65+ kg vs. ~6000+ kg) and hence the lack of necessity for extreme postural changes to maintain lower muscle stresses in order to maintain locomotor performance (Biewener, 1989, 1990). However, ostriches may also have a greater importance 706 for non-isometric muscle force-length properties in determining the limb orientation used 707 (Figures 6,7), as per the section above. Such speculations can be tested better once such 708 physiological data exist for ostrich muscles. Our data also do not strongly support Smith et 709 al.'s (2007) suggestion that hip extensor (or other muscle) moment arms are at peak values toward the end of stance phase (Figures 6-8). Overall, unfortunately the factors that 710 determine limb orientation in locomoting ostriches, as the largest extant striding biped (and 712 theropod dinosaur) available for study, remain inconclusive, leaving the application of such 713 principles to reconstructing limb orientations and locomotion in extinct theropods (e.g., 714 Hutchinson et al., 2005; Gatesy et al., 2009) on shakier empirical and theoretical ground.

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716 How accurate and repeatable are estimates of ostrich limb muscle moment arms?

717 Our Question 3 dealt with a methodological comparison among the three main studies 718 of ostrich pelvic limb muscle moment arms. Agreement seems fair overall, especially for 719 flexion/extension actions, but several main messages emerge from our comparisons, some of 720 which were also voiced by the other two studies of ostrich pelvic limb moment arms.

721 Circumstantial support for all three methods' accuracy also comes from tendon travel 722 measurements of cranial and caudal parts of the IL muscle in guineafowl by Carr et al. 723 (2011). General patterns (their Figure 7) for the IL moment arms about the knee (concave arc, 724 peaking $\sim 100^{\circ}$ knee angle in flexion) and the hip (increasing with extension) agree 725 reasonably well with these three ostrich studies (Figures 12, 16). However, all moment arms 726 for the ostrich IL muscle infer a switch to hip flexor action in strongly flexed poses, and little or no levelling off of the moment arm curve at strong hip flexion angles. 727

728 Key areas of disagreement between our results and those of B.A.S. and/or S.E.A. 729 include occasionally major differences in if, or how, muscles switch between flexion and extension (e.g., the AMB1 and AMB2, IC, ILFB about the hip; Figures 9-11), whether certain 730 731 muscles are flexors or extensors (e.g., the OM; see "Implications for ostrich limb muscle 732 function" below), or the absolute magnitudes or relative trends in the data (e.g., our nearconstant moment arms about the ankle for the FDL and gastrocnemius muscles; Figures 733 734 18,19; and for the digital muscles, Figure 20). We also found some differences in LAR and 735 ab/adduction moment arms about the hip for B.A.S.'s data, but these are likely explained by 736 differing muscle paths (e.g. via points and wrapping); see Figures 12-15. Bates and 737 Schachner (2012) acknowledge that their model could not use both via points and wrapping 738 surfaces for the same muscle, which explained the switch of their ILFB knee moment arm 739 from flexor to extensor with knee flexion (unlike Smith et al.'s (2007) data); our model only 740 exhibits this switch at extreme knee flexion ($\sim 150^{\circ}$ vs. 90° ; Figure 16). 741

Contrastingly, the "M. femorotibialis medialis" (see Appendix 1; equivalent to part of our FMTL; Figure 16, "FTE-Bates") muscle's moment arm changed with knee extension similar to other knee extensors in B.A.S.'s model, but S.E.A. found a progressive decline with knee extension. By comparing homologous muscles, it is evident that the experimental data ("FMTM-Smith" in Figure 16) match our model somewhat (FMTL; Figure 16), whereas the two distal parts of FMTL ("FTE" in Figure 16; B.A.S. and S.E.A. data) match each other somewhat, but only our model represents the anatomy and function of the actual medial head of M. femorotibialis (FMTM; Figure 16). Thus, a combination of incorrect classification of muscles (Appendix 1) and methodological differences explains divergence between some results. We noted similar problems with misidentifications (IFE, ITC, ITM muscles; Appendix 1) above in describing the results shown in Figure 10.

We raise the point here of the mistaken notion that tendon travel estimates of moment arms are free of errors in joint centre estimation, which is often repeated (e.g., Smith et al., 2007; Channon et al., 2010). This is only partly correct-- by using the virtual work principle 755 that moment arms are equal to the change in musculotendinous length ("tendon travel") per 756 unit of joint rotation, tendon travel experiments do minimize errors in estimating moment 757 arms, but still require joint centre estimates to calculate joint rotations, forming the traditional 758 x-axis of moment arm vs. joint angle plots. However, admittedly all studies carry this burden 759 of error; our model is different in using empirically measured 3D joint axes. Yet by attempting to restrict limb joints to flexion-extension axes, typical tendon travel experiments 760 still introduce another error, by introducing inevitable kinematic cross-talk between 761 762 flexion/extension movements and motions about other planes (i.e. limb joints that are 763 manually flexed/extended through a range of motion will also involve some motion in LAR 764 and ab/adduction-- see Rubenson et al., 2007). Only the most rigorously constrained studies, 765 which measure joint axes and constrain motions to strictly flexion-extension planes, avoid 766 this problem. Models like ours and B.A.S.'s can explicitly avoid it. Our model's data show 767 that, because moment arms of muscles covary with all joint angles (flex/extension, 768 ab/adduction, long-axis rotation) (Figures 12-16, S1-S4), changes in multiple joint angles at 769 once (i.e., kinematic cross-talk) will tend to produce different moment arms than changes in 770 one angle at a time.

771 Bates and Schachner (2012:p.1342) inferred that, because their model's outputs 772 matched experimental data, "predicted abduction/adduction and long-axis rotation moments 773 are good estimates". We find some important differences between our results and those of 774 either or both studies that somewhat weaken this inference. However, we reiterate and 775 celebrate that all three studies discussed here obtain broadly similar results for most muscles, 776 in particular emphasizing that moment arms are not constant for most muscles. Furthermore, both Smith et al. (2007) and Bates and Schachner (2012) noted limitations similar to thosementioned above.

779 Some comparative data exist from studies of other ratites or more restricted datasets 780 for ostriches. Hutchinson (2004a) provided estimates weighted by physiological cross-781 sectional area (A_{phys}) of "antigravity" (extensor) muscle moment arms for a 2D model of an ostrich, using the same specimen and hence congruent with our results. Alexander et al. 782 783 (1979) created a simple 2D geometric model of an ostrich whose limb dimensions were 784 similar to ours (cf. their Table I and our Table 1), obtaining comparable muscle moment arm 785 measurements: femorotibialis 5.1cm about the knee (ours ~5 cm), gastrocnemius 3.8 cm 786 about the ankle (ours ~4 cm), and digital flexors 3.2 cm about the MTP joint (ours ~2 cm). 787 This ostrich was 64% of the body mass of ours but had muscle masses about 85% of ours, 788 helping to explain its ~30% greater A_{phys} and F_{max} values (in addition, surely, to shorter 789 muscle fascicles; Equations 1, 2), and attributable to its wild-caught status as opposed to our 790 specimen's farm provenance. 791

Goetz et al. (2008) created a musculoskeletal model of an emu using a procedure grossly similar to ours, and obtained similar results—e.g. ILFB muscle having the largest hip extensor moment arm (but see Appendix 1). Troy et al. (2009) assumed that only the IFE, ITM, ITCR and PIFML muscles (homologous to ours) would resist hip adduction in their simplified emu model, but our analysis reveals that several more hip abductors exist, namely the IL, ILFB, FCM/L and CFP muscle groups (Table 4; Figures 14,15). See Lamas et al. (2014) for more consideration of emu muscle function.

There are anatomical and methodological reasons to suggest that musculoskeletal modelling approaches (this study and Bates & Schachner (2012)) can sometimes be more reliable than tendon-travel-based empirical data (e.g. Smith et al., 2007; also possibly Carr et al., 2011). First, the typical experimental method, as applied to date with birds, removes surrounding muscles and replaces whole 3D muscle bellies with quasi-2D strings (often lacking key "via points" or wrapping surfaces) that must alter the lines of action and thus moment arms of the *in situ* muscles. The potential for such alteration is experimentally testable, but existing studies of moment arms in various species (especially humans) already give strong precedent for this inference: analogous comparisons of "straight-line" 2D measurements (e.g., Jensen & Davy, 1975) vs. 3D medical imaging or in situ measurements (e.g., Young et al., 1993; Arnold et al., 2001) vs. anatomically-realistic 3D musculoskeletal models (Delp et al., 1999; Arnold et al., 2000; Kargo & Rome, 2002; Brown et al., 2003b; Burkholder & Nichols, 2004; Blemker et al., 2006; O'Neill et al., 2013) show better agreement between the latter two methods, and areas where simple 2D measurements are imprecise or misleading. To be fair, poorly designed musculoskeletal models can have the same problems—faithful representations of anatomical geometry are vital for both methods. Bates and Schachner (2012, their electronic supplementary material) note that their

815 model obtained an "extremely close match to the experimental data, particularly given the 816 level of intra-specific variation present in muscle moment arms for ostriches". Smith et al. 817 (2007) did attribute some of their experimental variation to individual differences in morphology that might alter moment arms, but as they noted some of the variation may have 818 819 instead been due to experimental error. Young et al. (1993), using a rigorous, validated in situ 820 experimental apparatus for muscles crossing cat ankle joints, still found some individual 821 variability that they attributed to muscles tethered by soft tissues as opposed to those restricted by bony wrapping surfaces or travelling in simple straight lines without any 822 823 obstacles. We caution that these different sources of error need to be weighed separately; 824 individual variability is an interesting biological reality, but experimental error needs minimizing. We suspect that some of the issues raised above about unrealistic representation 825 of musculoskeletal anatomy in tendon travel experiments might be due to experimental error. 826

Bates and Schachner (2012) also raised the latter issue, finding that trends evident in Smith et al.'s (2007) moment arm data would leave the hip extensors too weak to balance the hip joint moments during stance phase, with a premature shift to hip flexor action. Our results (Figs. 9-20) indicate that tendon travel measurements of moment arms often exhibit steeper changes with joint angle in some cases, or constant moment arms where models show good reason to suspect variable moment arms with joint angle.

833 O'Neill et al.'s (2013; also Howlowka & O'Neill, 2013) musculoskeletal modelling 834 and experimental analysis of chimpanzee pelvic limb moment arms came to similar conclusions as we have here, concerning model vs. tendon travel estimates of moment arms 835 (see also Brown et al., 2003a,b for horses). Importantly, they noted that non-linear trends in moment arm vs. joint angle data are to be expected, so tendon travel studies that enforce the fitting of largely linear (or simple polynomial) curves to their data may introduce errors. O'Neill et al. (2013) also cautioned that (p.3721) "the 3D orientation of the pelvis and thigh segments during tendon-excursion experiments can have a significant effect on moment arm metrics." Channon (2010) used cubic spline functions to characterize gibbon hindlimb muscle moment arms, presenting an improvement over the method used for ostriches by Smith et al. (2007) and considering some of the potential errors in tendon travel methods. Tightly controlled experiments with cats (Young et al., 1993) have shown good agreement with model-based moment arm data (Burkholder and Nichols, 2004).

Nonetheless, there is room for improvement in all methodologies. In particular, human experiments have shown that moment arms may depend on muscle activation levels (Maganaris, 2004; Tsaopoulos et al., 2007), a phenomenon that few moment arm studies have assimilated. Numerous studies have also shown that sarcomere or fibre length changes in real 3D muscles, as opposed to line segments, may be highly heterogeneous and complex (e.g., Blemker et al., 2005; Carr et al., 2011).

The accuracy of avian musculoskeletal models remains unresolved. Studies of moment arms for ostriches and other avian species badly need stronger validation tests – 854 ideally 3D, in vivo and across a range of behaviours - to test where different methods succeed 855 or fail. O'Neill et al. (2013) noted that model-based estimates of moment arms should be less 856 sensitive to the origins of muscles than to the insertions, and given that those insertions tend 857 to be more easily circumscribed in ostriches (as in chimpanzees), a focus on improving the 858 placement of muscle insertions could be powerful. At present, there is no published gold 859 standard measurement that the aforementioned studies can be compared against, and thus it is unclear which estimates of moment arms are truly better than others. Our judgements above 860 might prove to be incorrect. We assume here, except where noted, that our moment arm 861 862 estimates are generally an improvement over previous studies' because they are 3D, based on precise, subject-specific anatomical measurements of a single cadaver in situ, and incorporate 863 modern data on the 3D complexity of avian limb joint axes. However, our assumption of 864 865 improved accuracy demands a test against a gold standard, with clear criteria for what a "good" agreement between moment arm curves is; a question that no studies (including ours) 866 have answered. 867

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869 Model assumptions and potential refinements

Some simplifications of joint systems were necessary in our model but could be
improved with later iterations. The tibio-fibular articulation is slightly mobile in ostriches
(Fuss, 1996) and other birds, but we maintained it as an immobile joint. Likewise, the
(proximal; see Regnault et al., 2014) patella surely translates during knee flexion/extension in
birds as in humans (e.g. Walker et al., 1988) but we maintained it in the same resting position
(with respect to the femur), represented simply by a wrapping surface. Adding such
translation would influence the moment arm curves for knee extensor muscles. The intertarsal

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877 (ankle) joint's motions during swing phase (extreme dorsiflexion) seemed unrealistic, 878 laterally rotating the tarsometatarsus to a seemingly disarticulated position, but we kept this as-is in the model rather than invent a subjective solution, as it would have minimal influence 879 880 on our results here and maintained strict fidelity to our anatomical and kinematic data. Future implementations of the model emphasizing ankle joint mechanics may need to adjust these 881 kinematics. The proximal interphalangeal joint of digit III was kept immobile in the model as 882 883 our kinematic data lacked its joint motion, but the model has the capacity to allow the joint to 884 flex and extend if desired (Table 1), and could involve internal mechanisms such as those 885 described by Schaller et al. (2011) if necessary for research questions addressed with it.

Our model's muscles were simplified, as the Methods and Supplementary Text explain. Our digitizing procedure, performed in 2002, was simplistic (similar to that of Burkholder and Nichols, 2004), whereas more recent techniques have fused CT and MRI imaging modalities to produce quite accurate and complex 3D musculoskeletal models (e.g., Zarucco et al., 2005, Harrison et al., 2014). Real muscles have complex 3D structure but we have simplified them into basic Hill model muscles of 2D structure. Internal tendons were observed in some muscles (e.g., M. iliotrochantericus caudalis, Mm. gastrocnemii, many digital flexors; Gangl et al., 2004). The Hill model does not discretely represent these features, which can affect muscle forces and gearing. Ligaments and other passive tissues were not represented in our model, and these would be particularly important features to consider in a complete dynamic model, as Haughton (1864) suggested and Schaller et al. (2009) demonstrated experimentally. Finally, a test of the validity of our moment arm estimates against a "gold standard" empirical measurement (see above) with explicit criteria for an acceptable level of precision would be immensely valuable.

Yet like any model, simple or complex, our model is a useful starting point for a continually iterative process of improvement that progressively approximates reality. It has the advantage of being able to estimate muscle forces, moments, and length changes that are unlikely or even impossible to be measurable *in vivo*, especially simultaneously across a full stride.

906 Implications for ostrich limb muscle function

907 Table 4 shows our classification of the major actions of ostrich pelvic limb muscles 908 (also see Lamas et al., 2014 for emu muscles). This approach, as opposed to the classical 909 perspective in comparative anatomy and biomechanics, emphasizes the three-dimensional 910 nature of avian limbs and the capacity of muscles to generate moments about many joints and 911 degrees of freedom (see also the categorization for cat ankle muscles provided by Young et 912 al. (1993); also model-based assessments using isometric force-fields by Kargo & Rome 913 (2002) and endpoint forces by Burkholder & Nichols (2004)). Standard functional anatomy 914 papers tend to emphasize flexion/extension and only present other actions (or any actions 915 about other joints than those deemed to be the main joints of action) in a seemingly arbitrary 916 fashion (e.g., Smith et al., 2007: Table 2; cf. our Table 4), with at least the implicit 917 assumption that non-parasagittal actions are less important. Our approach also reveals how 918 the functions of some muscles have been misclassified before (or remain uncertain), for 919 ostriches and perhaps for other avian species--to the degree that ostriches are representative 920 of ancestral muscle functions, which remains to be determined. The degree of uncertainty 921 about avian pelvic limb muscle function particularly applies to 3D actions about the hip joint; many muscles have actions (e.g., flexion/extension) that switch depending on the joint angles 922 923 adopted.

Examples of somewhat surprising 3D actions of hip muscles include the M.
obturatorius medialis (OM), which we find to mainly fulfill a lateral rotator and flexor action
at the hip, whereas Smith et al. (2006,2007), Bates and Schachner (2012), Bates et al.

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927 (2012a,b) favour an extensor action of this muscle (in addition to lateral rotation) in ostriches 928 and other archosaurs. In the model of B.A.S., this seems to be at least partly due to a distal 929 insertion of the OM muscle on the femur, rather than very proximally onto the trochanteric 930 crest (Gangl et al., 2004; Zinoviev, 2005). Furthermore, some "antigravity muscles" act in 931 one direction synergistically with other muscles, resisting gravity, but in other directions they 932 act antagonistically to gravity. Clear examples include the PIFML and CFP muscles, which 933 are presumably active in hip extension but incur lateral rotator and abductor moments about 934 the hip (Table 4). These muscles probably oppose other antigravity muscles such as the ITC in medial rotation and potential adductors such as AMB1 (cf. moment data in Rubenson et 935 936 al., 2011).

The analysis of Rubenson et al. (2011:Figure 7) indicates that hip adduction, not abduction, must be resisted during stance phase in locomoting ostriches, and thus abductor muscle activity is predicted, a moment that many hip extensors create anyway. However, hip adduction capacity is far more limited -- only the IC, AMB1, 2 and IFI muscles have clear actions in hip adduction. Smith et al. (2006:Table 2) assigned adductor actions to other muscles such as the flexor cruris (FC) heads, PIFML and OM whereas we find these to be abductors. Indeed, the actions of the two heads of AMB may oppose each other (Table 4), so it would be interesting to know how they are coordinated. The ITC muscle's parts have clear roles in medial rotation, but their actions in flexion/extension vary with limb posture (see also Gatesy, 1994), rendering it less clear whether (or when) they play a predominant hip extensor (e.g. Rubenson et al., 2006) or hip flexor (e.g. Smith et al., 2006,2007) role in ostriches or other birds. How any birds balance this complex interaction of long-axis and ab/adductor moments at the hip or other joints remains almost unexplored (but see Gatesy 1994), yet modelling (and simulation) approaches such as ours offer one way to tease apart the complexity. Bates and Schachner (2012) found that Alligator and Struthio had similar hip extensor moment arms but there were large abduction and small adduction moment arms in their ostrich, along with large long-axis rotation moment arms. The functional and evolutionary implications of these differences remain unclear, and dependent on understanding force balance about the hip joint in extant archosaurs such as Struthio.

956 Complex function is not restricted to proximal muscles, however. Complex anatomy 957 of distal limb muscles is a pernicious problem in avian locomotor biomechanics, and difficult 958 to render realistically in biomechanical models such as ours. As the Supplementary Text 959 describes, we could not model all origins (or subdivided tendons; e.g., digital flexors) of all 960 muscle parts. Indeed, in some cases the origins are diffuse-e.g. M. gastrocnemius medialis 961 originates mainly from the medial side of the proximal tibiotarsus, but the surrounding fascia 962 it is attached to continue proximally past the knee joint, via the proximal patella and other 963 structures. It is not clear if some of these distal muscles exert important moments about the 964 knee joint (some forces may be going directly to their distal origins from the tibiotarsus), and 965 the dynamics of the patella (not represented in our model except as a static wrapping surface) 966 further complicates matters. Thus it is unclear how forces are balanced across ostrich (or 967 other avian) knees, complicating comparisons with other species (e.g. Higham et al., 2008; 968 Andrada et al., 2013).

969 Young et al. (1993) and Johnson et al. (2012) noted that some muscles in cat and rat 970 hindlimbs seemed to have intrinsically stabilizing properties, shifting from flexor to extensor 971 moment arms in a linear fashion with increasing joint flexion. Eight modelled ostrich limb 972 muscles also show this pattern: the AMB1, AMB2, IC, ITCa, ITCp, ITM, ITCR and ISF 973 exhibit stabilization function in flexion-extension (Figures 9,10). Weaker evidence for self-974 stabilization is present for the OM muscle in hip ab/adduction (Figure 14) and the four ankle 975 flexors in flexion/extension (TCf, TCt, EDL, and FL; Figure 18), so any self-stabilization properties must be interpreted as being largely restricted to the hip's flexion-extension 976

function (see also Table 4). Judging from these hip muscles' paths, their long lines of action
(due to the elongate pelvis and limb) running close to the hip seem to predispose them to
these intrinsic stabilization properties. These patterns deserve more examination in a
comparative context with other species in the future -- for example, similar hip muscles show
similar moment arm patterns that hint at intrinsic stabilization in various extinct dinosaurs
and other archosaurian reptiles (Hutchinson et al., 2005,2008; Bates and Schachner, 2012;
Bates et al, 2012a,b; Maidment et al., 2013).

985 Musculoskeletal models of limb function: past, present and future
 986 A wide variety of studies have used musculoskeletal mode

A wide variety of studies have used musculoskeletal models to reconstruct limb function in extant and extinct animals, but there remains little agreement for standards of model design, analysis and validation. The same software (SIMM) or other packages (GaitSym, Anybody, varieties of Adams, etc.) has been used to estimate limb muscle moment arms in other extant species including chimpanzees and other hominins (O'Neill et al., 2013; Holowka & O'Neill, 2013; and references therein), horses (Brown et al., 2003a,b; Zarucco et al., 2006; Harrison et al., 2010), domestic cats (Burkholder & Nichols, 2004), rats (Johnson et al., 2008), emus (Goetz et al., 2008), Alligator and ostriches (Bates and Schachner, 2012), frogs (Kargo and Rome, 2002; Kargo et al., 2002), cockroaches (Full & Ahn, 1995), and others, in addition to simpler past approaches (e.g. for small mammals, McClearn, 1985). It is not clear which software packages most accurately estimate muscle moment arms, but our comparisons with the data from Bates and Schachner (2012) suggest some advantages of our approach. Regardless, comparative biologists seeking to harness the power of musculoskeletal modelling techniques have much to learn from the considerable progress made in analyzing and validating similar models of human limb muscles (e.g., Delp et al., 1990,1992,1999; Arnold et al., 2000,2001; Holzbauer et al., 2005; Nikooyan et al., 2011).

There has been a recent flourishing of musculoskeletal models of extinct taxa, especially hominins (e.g., Australopithecus in Nagano et al., 2005) and dinosaurs or other archosaurs. Theropod dinosaurs have featured prominently, including Allosaurus, 1004 Struthiomimus (Bates and Schachner, 2012; Bates et al. 2012a), Tyrannosaurus (Hutchinson 1005 1006 et al., 2005), and Velociraptor (Hutchinson et al., 2008), among other taxa (Allen, 2010), and 1007 more recently sauropodomorph (Mallison, 2010a,b) as well as ornithischian dinosaurs have 1008 enjoyed a new focus (Mallison, 2010c; Bates et al., 2012b; Fujiwara and Hutchinson, 2012; 1009 Maidment et al., 2013). Other extinct taxa studied have included the stem crocodile Poposaurus (Bates and Schachner, 2012) and related taxa (Molnar, 2014), a pterosaur (Costa 1010 1011 et al., 2013), and the early tetrapod Ichthyostega (Pierce et al., 2012). Whether the focus is on 1012 limb joint ranges of motion, muscle moment arms, or dynamic simulation, there is a need for 1013 more scrutiny of models of extant taxa to establish how accurate and reliable these modelling 1014 approaches truly are, and ultimately a renewed examination and synthesis of those findings 1015 with past studies of extinct taxa. Studies using models to estimate muscle moment arms need 1016 to consider not only this, but also how or whether those parameters actually matter for 1017 particular muscles, joints, behaviors or species (see also Bates and Schachner, 2012; Bates et 1018 al., 2012b; Maidment et al., 2013). To the degree that general principles of moment arm 1019 usage exist, they should clarify under what circumstances a moment arm is optimized to 1020 perform a certain function. An understanding of this link between muscle mechanics, 1021 kinematics and moment arms could link the disparate palaeobiological/comparative studies 1022 on limb joint ranges of motion and orientations (e.g., Mallison, 2010a,b,c; Pierce et al., 2012; 1023 Molnar, 2014) with those of limb muscle moment arms (Hutchinson et al., 2005,2008; Allen, 1024 2010; Bates and Schachner, 2012; Bates et al., 2012b; Fujiwara and Hutchinson, 2012; Maidment et al., 2013). Such a synthesis could lead to a robust, sustainable future for 1025

1026 comparative musculoskeletal modeling and simulation, and new insights into the evolution of 1027 musculoskeletal function and locomotor dynamics.

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APPENDIX 1:

A few observations from our dissections revealed differences in muscle identifications from previous studies, which themselves have had much disagreement (reviewed in Gangl et al., 2004; Zinoviev, 2006). However, generally our findings agree well with the excellent study by Gangl et al. (2004). Zinoviev (2006) made some amendments to the latter study that our dissections independently have confirmed.

First, rather than treat "M. pectineus" as a novel muscle unrelated to M. ambiens (e.g., Mellett, 1994; Gangl et al., 2004) or as a part of M. femorotibialis medialis (= FMTM or "internus"; Zinoviev, 2006), we refer to the muscle here as M. ambiens 1 (AMB1; for its pubic origin as in most other Reptilia; Hutchinson, 2002) and refer to the muscle with the derived iliac origin as M. ambiens 2 (AMB2; i.e., the dorsal head) (Table 2). We do not intend for this to be a formal nomenclatural change but it is used for convenience within this paper and to suggest that the homologies (and thus formal nomenclature) of these two muscle heads deserve reconsideration. Our nomenclature avoids confusion with the mammalian M. pectineus which has no homology with these muscles, and maintains the same nomenclature for the homologous M. ambiens (whether one or two parts) in extant Sauropsida. We deem these homologies and nomenclature marginally more parsimonious than other interpretations (reviewed by Zinoviev, 2006), the latest of which (favored by Zinoviev, 2006) requires loss of the origin of M. ambiens from the pectineal (preacetabular) process that is plesiomorphically present in birds (Hutchinson, 2002) and gain of a new head of the FMTM 1059 muscle (the "pars pectineus"). In our scenario, mainly a new head of M. ambiens (AMB2) is 1060 required; a phenomenon that is not unknown in other birds and is pervasive in Crocodylia (Hutchinson, 2002).

1062 Second, we agree with Zinoviev (2006) that Gangl et al. (2004) confused the two 1063 crura of "M. iliofemoralis externus" with M. iliotrochantericus caudalis, which engendered further errors in identifying other muscles. We have confirmed this from dissections of three 1064 1065 additional ostrich cadavers, and hence the position of our M. iliotrochantericus caudalis 1066 (ITCa,ITCp in Table 2) differs from theirs (also data in Smith et al., 2006,2007). We consider 1067 M. iliofemoralis externus (IFE) to be weakly differentiated from the latter muscle, also 1068 intimated by Gangl et al. (2004:p.113; and Gadow, 1880), but we represent it in our model as 1069 a separate head (IFE), again agreeing with Zinoviev (2006). This is corroborated by the main muscle in question (ITC) having a large internal tendon, preacetabular position (between the 1070 1071 heads of Mm. iliotibiales; IC and IL), and overlaying Mm. iliotrochanterici medius et 1072 cranialis (ITM, ITCR), which are traits diagnostic of M. iliotrochantericus caudalis, not "M. iliofemoralis externus" as in Gangl et al. (2004). This then explains why the latter study did 1073 1074 not find M. iliotrochantericus medius - it identified it as M. iliotrochantericus caudalis, 1075 noting some fusion and distolateral insertion with M. iliotrochantericus cranialis, which are

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diagnostic of M. iliotrochantericus medius instead. Hence our M. iliotrochantericus medius
(ITM) corresponds to their "M. iliotrochantericus caudalis"; but our M. iliotrochantericus
cranialis (ITCR) corresponds to theirs. Smith et al. (2006,2006) perpetuated the errors caused
by Gangl et al.'s misidentifications, which complicates comparisons with our data (see our
Results section, Fig. 10). Bates and Schachner (2012) appear to have avoided these errors.

A third point of discordance between this study and Gangl et al.'s (2004) is that we 1081 1082 consider the latter study's "Mm. femorotibiales externus et medius" to be two parts 1083 (superficial and deep) of M. femorotibialis lateralis (FMTL; vide Zinoviev, 2006), because 1084 this avian muscle typically originates on the lateral surface of the femur, deep to M. 1085 iliotibialis lateralis (IL), as the former two parts do. Gangl et al.'s (2004) "Mm. 1086 femorotibiales internus et accessorius" then correspond to our Mm. femorotibiales medialis et intermedius (FMTM, FMTIM), because their topological connections more closely match 1087 these muscles in birds. Zinoviev (2006) again gave a detailed correction that we concur with, 1088 whereas Smith et al. (2006,2007) and possibly Bates and Schachner (2012) used Gangl et 1089 al.'s (2004) misidentifications.

Accepting these identifications renders ostrich anatomy more similar to other birds: M. iliotrochantericus caudalis remains large and originating dorsal to M. iliotrochantericus medius, which originates cranial to M. iliotrochantericus cranialis, and the insertions of these muscles retain similar proximodistal positions (the first on the lateral surface of the proximal femur, the second two sequentially distal on the craniolateral edge of the trochanteric crest; Rowe, 1986; Hutchinson, 2002). Furthermore, Mm. femorotibiales in our view then match avian anatomical positions more reasonably.

However, ostrich pelvic muscle anatomy is still derived in the large relative size of M. iliofemoralis internus (IFI; "cuppedicus" of Rowe, 1986; Table 2) and the two distinct heads of M. ambiens (i.e., the novel dorsal head, AMB2). Additionally, M. iliotrochantericus medius (ITM) is slightly derived in having a markedly more dorsal position (required by the dorsally shifted M. ambiens and M. iliofemoralis internus (IFI)) than in typical neornithines, and Mm. femorotibiales have strong division of the lateral head (FMTL) into two parts, but not a novel "accessorius" head. Furthermore, M. obturatorius medialis (OM) in ostriches is extraordinary in its mostly lateral origin, whereas normally in birds it is confined to the inside of the pelvis.

1107 Similar problems with muscle identifications vex other studies of ratite limb form and function. Patak & Baldwin's (1998) anatomical description of emu pelvic limb muscles was 1108 1109 used to formulate Goetz et al.'s (2008) musculoskeletal model of an emu, incorporating 1110 analogous errors that deserve correction in future studies (see Lamas et al., 2014). The issues 1111 outlined in this appendix drive home the point that a firm grasp of avian myology, ideally based upon understanding of fundamental groups of muscles likely present in the ancestral 1112 1113 neornithine bird and hence homologous among its living descendants, is critical for any 1114 biomechanical and comparative analyses of muscle form, function and evolution (see also 1115 Zinoviev, 2006).

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1552 Figure Captions

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Figure 1. Digitizing apparatus used during anatomical dissection of ostrich. "LED Ref"
indicates the proximal (in trochanteric crest of the femur) and distal (in tibiotarsus by the
ankle) reference frames, "Dig. Probe" indicates the digitizing probe used to collect
landmarks.

Figure 2. Ostrich model joint axes (x, y, z) shown in right lateral (A) and oblique right
dorsolateral (B) views. The x-axis corresponded to ab/adduction, the y-axis to long-axis
rotation, and the z-axis to flexion/extension.

Figure 3. Ostrich musculoskeletal model in right lateral view, with muscle-tendon units labeled (red lines). See Table 2 for muscle abbreviations.

Figure 4. Ostrich musculoskeletal model in right caudolateral view, with muscle-tendon units labeled (red lines). See Table 2 for muscle abbreviations.

Figure 5. Ostrich musculoskeletal model: wrapping surface examples. See Table 2 for muscle abbreviations. Lateral (A), caudolateral (B), and craniolateral (C) views of eight muscle wrapping objects (in blue), as half and whole cylinders, ellipses and a torus. The PIFML and ILFB wrapping surfaces are shown as meshes, for added clarity.

Figure 6. Maximal muscle moments about proximal limb joints (hip and knee), for representative walking and running trials (see Methods). "F-L" curves incorporate effects of muscle force-length properties into moment calculations; "Fmax" curves only assume maximal isometric muscle stress and thus ignore F-L effects.

Figure 7. Maximal muscle moments about distal limb joints (ankle and metatarsophalangeal [MTP]), for representative walking and running trials (see Methods). See caption for Figure 6.

Figure 8. Sum of extensor moment arms (left column) or flexor moment arms (right column) normalized by sum of maximal extensor or flexor moment arms, plotted against extension or flexion joint angle for the hip, knee and ankle joints (MTP joint data follow Figure 20), with representative mid-stance limb poses for walking and running indicated.

Figure 9. Hip flexor/extensor moment arms plotted against joint angle for key proximal thigh muscles in our model, with corresponding data from Smith et al. (2005) labelled as "Smith" and from Bates and Schachner (2012) labelled as "Bates". Extreme extended/flexed right hip joint poses shown along the x-axis. Muscle abbreviations are in Table 2.

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Figure 10. Hip flexor/extensor moment arms plotted against joint angle for key proximalthigh muscles. See caption for Figure 9.

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Figure 11. Hip flexor/extensor moment arms plotted against joint angle for key proximal
thigh muscles. See caption for Figure 9.

1599 Figure 12. Hip long-axis rotation (LAR) moment arms plotted against hip flexion/extension

1600 joint angle for key proximal thigh muscles. See caption for Figure 9.

- 1607 1608 1609 1610 1611 1612 1613 1614 1615 1616 1617 1618 1619 1620 1621 1622 1623 **J**1624 1625 **D**1626 1627 1628 1629 1630 1631 1632 1633 1634 1635 1636
- 1602 Figure 13. Hip long-axis rotation (LAR) moment arms plotted against hip flexion/extension 1603 joint angle for key proximal thigh muscles. See caption for Figure 9. 1604
- Figure 14. Hip abduction/adduction moment arms plotted against hip flexion/extension joint 1605 angle for key proximal thigh muscles. See caption for Figure 9. 1606
 - Figure 15. Hip abduction/adduction moment arms plotted against hip flexion/extension joint angle for key proximal thigh muscles. See caption for Figure 9.
 - Figure 16. Knee flexor/extensor moment arms plotted against knee flexion/extension joint angle for key thigh muscles. See caption for Figure 9.
 - Figure 17. Knee flexor/extensor moment arms plotted against knee flexion/extension joint angle for key thigh and distal knee muscles. See caption for Figure 9.
 - Figure 18. Ankle flexor/extensor moment arms plotted against ankle flexion/extension joint angle for key muscles crossing the ankle. See caption for Figure 9.
 - Figure 19. Ankle flexor/extensor moment arms plotted against ankle flexion/extension joint angle for the M. gastrocnemius muscle group. See caption for Figure 9.
 - Figure 20. Metatarsophalangeal (MTP) joint flexor/extensor moment arms plotted against MTP flexion/extension joint angle for digital flexors (left) and extensors, plus tendinous connection of M. fibularis longus (right). See caption for Figure 9.

Supplementary files:

Supplementary Text

Supplementary Figures S1-S4: Hip muscle moment arms in long-axis rotation (LAR) or ab/adduction plotted against hip LAR or ab/adduction angles (cf. Figures 12-15 plotted against hip flexion/extension angles), for key proximal thigh muscles. See caption for Figure 9.

1637 Supplementary movie:

1638 1639 Movie S1: Musculoskeletal model of the right and left pelvic limbs of an ostrich, visualized 1640 statically to show 3D anatomy represented in the model; posed at mid-stance of running 1641 (right limb) and corresponding swing phase (left limb).

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Figure 1(on next page)

Figures 1-20

Captions in main text file (end)



Figure 1









Figure 3



Figure 4

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Figure 5



Figure 6









Figure 8



Figure 9



Figure 10



Figure 11



Figure 12



Figure 13



Figure 14



Figure 15



Figure 16



Figure 17



Figure 18





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Tables (all)

All tables with captions

Tables

	centre	centre	centre	Motion	
Joint or segment	x (m)	y (m)	z (m)	axes	Ranges of motion (°)
Pelvis	0	0	0	x,y,z	[-180/180; -180/180; -180/180]
Hip (acetabular/antitrochanteric)	0	0	0.0355	x,y,z	[-45/45; -45/45; -65/10]
Knee (femorotibial)	0	-0.2338	0.0543	x,y,z	[-45/45; -45/45; -180/10]
Ankle (intertarsal)	0	-0.442	0	X,Z	[-45/45; -10; -10/180]
Metatarsophalangeal (MTP III)	0	-0.426	0	Z	[5; 24; -180/90]
[Proximal interphalangeal (D III)]	[0	-0.089	0]	[Z]	[not estimated]

Table 1. Joint axes for the ostrich musculoskeletal model. Each joint centre is listed in (x,y,z)-coordinate space as a distance from the segment origin. The pes was 0.141m long and an interphalangeal joint's location is noted here in the final row, but was not included in the model. Each joint was defined relative to the one proximal to it, with the pelvis segment placed at the origin of the world coordinate system.

		Muscle			Maximal
		mass;	Fascicle	Pennation	isometric
Muscle		m _{musc}	length;	angle; θ	force;
abbreviation	Muscle full name	(kg)	L (m)	(°)	$F_{max}(N)$
IC	M. iliotibialis cranialis	0.3788	0.174	0	615
ILa	M. iliotibialis lateralis (cranial part)	1.074	0.174	0	875
ILp	M. iliotibialis lateralis (caudal part)		0.174	0	875
AMB1	M. ambiens, ventral (pubic) head	0.093	0.039	10	672
AMB2	M. ambiens, dorsal (iliac) head	0.1994	0.044	15	1240
FMTL	M. femorotibialis lateralis	0.3181	0.088	15	992
FMTIM	M. femorotibialis intermedius	0.387	0.084	25	1180
FMTM	M. femorotibialis medialis	0.272	0.089	30	753
ILFBa	M. iliofibularis (cranial part)	1.0623	0.176	0	867
ILFBp	M. iliofibularis (caudal part)		0.176	0	867
ITCa	M. iliotrochantericus caudalis (cranial part)	0.3114	0.064	25	622
ITCp	M. iliotrochantericus caudalis (caudal part)		0.064	25	622
IFE	M. iliofemoralis externus	0.03264	0.025	25	331
ITM	M. iliotrochantericus medius	0.0256	0.058	0	125
ITCR	M. iliotrochantericus cranialis	0.0432	0.053	10	228
IFI	M. iliofemoralis internus	0.0407	0.041	0	284
FCM	M. flexor cruris medialis	0.1192	0.036	35	767
FCLP	M. flexor cruris lateralis pars pelvica	0.3182	0.24	0	376
FCLA	M. flexor cruris lateralis pars accessoria	0.0211	0.125	0	47.8
ISF	M. ischiofemoralis	0.0348	0.033	15	290
PIFML	Mm. puboischiofemorales medialis + lateralis	0.1273	0.089	15	389
OM	M. obturatorius medialis	0.457	0.055	25	2160
CFP	M. caudofemoralis pars pelvica (et caudalis)	0.3069	0.108	15	778
GL	M. gastrocnemius pars lateralis	0.5706	0.12	20	1269
GIM	M. gastrocnemius pars intermedia	0.2526	0.125	15	552
GM	M. gastrocnemius pars medialis	0.762	0.094	20	2160
FL	M. fibularis longus	0.4791	0.081	20	1570
FDL	M. flexor digitorum longus	0.1424	0.048	20	782
FPPD3	M. flexor perforans et perforatus digitorum 3	0.0822	0.025	30	798
FPD3	M. flexor perforans digitorum 3	0.1605	0.017	35	2220
FPD4	M. flexor perforans digitorum 4	0.0955	0.026	20	992
FHL	M. flexor hallucis longus	0.0505	0.04	25	324
EDL	M. extensor digitorum longus	0.115	0.049	30	576
TCf	M. tibialis cranialis (femoral head)	0.165	0.045	25	474
TCt	M. tibialis cranialis (tibial head)		0.045	25	474

Table 2. Muscles included in the ostrich musculoskeletal model, with their associated abbreviations and physiological/architectural parameters. Data were obtained via dissection. Blank cells for muscle masses (ILp, ILFBp, ITCp, TCt) indicate that the second part of the muscle shares the mass value, which was divided equally to calculate A_{phys} and hence F_{max} .

Muscle(s)	Location	Shape	r (x)	r (y)	r (z)	t (x)	t (y)	t (z)	Radius	Length	_
ILFB	pelvis	cylinder	17.11	57.87	-34.76	0.0309	-0.0609	0.0622	0.095	1.000	
PIFML	pelvis	cylinder	0.92	-5.72	-29.06	0.0020	0.0820	0.1000	0.170	0.500	
CFP	pelvis	cylinder	-8.51	0.41	35.68	-0.0211	0.0722	0.1396	0.090	0.500	
ITC,ITM	femur	cylinder	-16.00	19.60	0.00	0.0086	-0.0017	-0.0067	0.020	0.500	
ITCR	femur	cylinder	-16.00	-2.17	-0.19	0.0395	-0.0379	-0.0034	0.020	0.500	
ITCR	femur	cylinder	74.00	0.00	-19.60	0.0086	-0.0882	0.0181	0.055	0.500	
GL,GIM,FHL,FPD3	femur	cylinder	37.16	40.49	-20.39	-0.0197	-0.2006	0.0799	0.015	0.200	
ILFBp	femur 🕖	torus	51.43	-21.08	-22.24	-0.0202	-0.1827	0.0609	0.01*	0.08*	
FMTIM,FMTL	tibiotarsus	cylinder	0.00	0.00	44.69	-0.0014	0.0103	0.0093	0.038	0.500	
GL,GIM,GM	tibiotarsus	cylinder	0.00	0.00	0.00	-0.0058	-0.4435	-0.0090	0.040	0.150	
FP&PD3,FPD3,FPD4	tibiotarsus	cylinder	6.47	-7.64	40.43	0.0031	-0.4537	0.0090	0.030	0.200	
FDL,FHL	tibiotarsus	cylinder	0.00	0.00	40.00	-0.0014	-0.4501	0.0001	0.030	0.500	
AMB2	tibiotarsus	cylinder	0.00	0.00	0.00	0.0250	0.0126	-0.0012	0.038	0.100	
IC	tibiotarsus	cylinder	0.00	5.00	0.00	-0.0143	0.0098	-0.0054	0.030	0.500	
EDL,TCf,TCt	tibiota <mark>rsu</mark> s	cylinder	3.37	-15.47	0.32	0.0111	-0.4530	0.0009	0.020	0.100	
FP&PD3,FPD3,FL	tarsometatarsus	cylinder	0.64	-6.17	-1.51	-0.0020	-0.4296	0.0002	0.023	0.100	
FDL,FHL	tarsometatarsus	cylinder	-3.48	-13.61	-0.29	0.0025	-0.4319	-0.0024	0.022	0.100	
FPD4	tarso <mark>meta</mark> tarsus	cylinder	0.00	-35.00	0.00	0.0002	-0.4326	0.0025	0.017	0.100	
EDL	tarsometatarsus	cylinder	-1.19	-2.69	0.68	0.0029	-0.4312	-0.0152	0.023	0.100	
Musala	Location	Shana	r (v)	r (11)	r (7)	$t(\mathbf{v})$	$t(\mathbf{x})$	$t(\mathbf{z})$	Radius	Radius	Radius
IVIUSCIC II	<u>noluis</u>	allingei	$\frac{1(\lambda)}{2.70}$	<u>1 (y)</u> 1 44	<u>1 (2)</u> 100 02	$\frac{L(X)}{0.0020}$	<u>(y)</u>	<u>1 (2)</u> 0 06 40	<u>(A)</u> 0.100	<u>(y)</u>	<u>(Z)</u>
	pervis	empsoid	-2.79	-1.44	-100.93	-0.0020	-0.0020	0.0049	0.100	0.200	0.05
IFE	pelvis	ellipsoid	0.00	0.00	0.00	-0.0010	0.0110	0.0302	0.090	0.060	0.06
FCLP	pelvis	ellipsoid	-17.65	8.20	-84.13	-0.2258	-0.0290	0.0532	0.075	0.500	0.050

Table 3. Muscle wrapping surfaces assumed in the ostrich musculoskeletal model, with dimensions. Examples are in Figure 5.

Musele	Action Hip F/E	Hip LAR	Hip Ab/Ad	Knee F/F	Ankle F/F	MTP F/F
IC	F+*	M	AD+	F/E	1,2	1,12
ILa	F/E	M/L	AB+	E+		
ILp	E+	M/L	AB+	E+		
AMB1	E*	L	AD	F		
AMB2	\mathbf{F}^*	M/L	AD	E		
FMTL	-			E+		
FMTIM				Е		
FMTM				F		
ILFBa	Е	М	AB	F+		
ILFBp	E+	М	AB	F+		
ITCa	F/E*	M+	AB/AD			
ITCp	F/E [*]	M+	AB/AD			
IFE	F	M/L	AB			
ITM	F/E*	М	AB/AD			
ITCR	F/E [*]	M+	AB/AD			
IFI	F	M/L	AD			
FCM	Е	М	AB	F		
FCLP	E+	M+	AB+	F		
FCLA	Е	М	AB			
ISF	F/E*	L	AB			
PIFML	Е	L	AB			
OM	F+	L+	AB/AD*			
CFP	Е	L	AB			
GL				F	E+	
GIM				F	Е	
GM				(F/E)	E+	
FL				Е	F^{*}	F
FDL					E+	F+
FPPD3				(F/E)	E+	F+
FPD3				(F/E)	E+	F+
FPD4				(F)	E+	F+
FHL				(F)	Е	F
EDL					$F+^*$	E+
TCf				0	$F+^*$	E+
TCt					$F+^*$	E+

Table 4. Muscle actions, following results from Figures 9-20, to describe the major 3D potential functions of each ostrich pelvic limb muscle. Classifications: E=extensor, F=flexor, M=medial (internal) rotator, L=lateral (external) rotator, AB=abductor, D=adductor, 0=no moment arm *per se* despite crossing the joint. Blank cells indicate the muscle does not cross or act about the joint. "+" signs added to classifications indicate a major potential role in these functions based upon moment arm and muscle relative size (i.e., moment generation capacity), subjectively assessed. "/" combinations (F/E; M/L; AB/AD) indicate a strong sensitivity of muscle moment arm, and hence action, to joint angle. Annotation with an asterisk indicates a potential role for intrinsic stabilization about that axis of motion (see Discussion). "()" indicates that our model's single origin for each muscle (or part thereof) did

not allow such an action, but sub-parts of those muscles might have such actions if modelled in more detail.

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