# **RVC OPEN ACCESS REPOSITORY – COPYRIGHT NOTICE**

This author's accepted manuscript may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Self-Archiving.

The full details of the published version of the article are as follows:

TITLE: Relating neuromuscular control to functional anatomy of limb muscles in extant archosaurs

AUTHORS: Cuff, A R; Daley, M A; Michel, K B; Allen, V R; Lamas, L P; Adami, C; Monticelli, P; Pelligand, L; Hutchinson, J R

JOURNAL: Journal of Morphology

PUBLISHER: Wiley

PUBLICATION DATE: 8 March 2019 (online)

DOI: https://doi.org/10.1002/jmor.20973



1 2	Relating neuromuscular control to functional anatomy of limb muscles in extant archosaurs
3	Short title: Archosaur EMG patterns
4 5	Andrew R. Cuff <sup>1</sup> , Monica A. Daley <sup>1</sup> , Krijn B. Michel <sup>1</sup> , Vivian R. Allen <sup>1</sup> , Luis Pardon Lamas <sup>1,2</sup> , Chiara Adami <sup>3</sup> , Paolo Monticelli <sup>3</sup> , Ludo Pelligand <sup>3</sup> , John R. Hutchinson <sup>1</sup>
6 7	<sup>1</sup> Structure and Motion Laboratory, Department of Comparative Biomedical Sciences, Royal Veterinary College, Hawkshead Lane, North Mymms, Hertfordshire, AL9 7TA, United Kingdom.
8 9	<sup>2</sup> Current address: Faculdade de Medicina Veterinária, Universidade de Lisboa, Pólo 7, Universitário da Ajuda, 1300-477, Lisboa, Portugal.
10 11	<sup>3</sup> Queen Mother Hospital, Department of Clinical Science and Services, Royal Veterinary College, Hawkshead Lane, North Mymms, Hertfordshire, AL9 7TA, United Kingdom.
12	
13	Abstract
14	Electromyography (EMG) is used to understand muscle activity patterns in animals. Understanding
15	how much variation exists in muscle activity patterns in homologous muscles across animal clades
16	during similar behaviours is important for evaluating the evolution of muscle functions and
17	neuromuscular control. We compared muscle activity across a range of archosaurian species and
18	appendicular muscles, including how these EMG patterns varied across ontogeny and phylogeny, to
19	reconstruct the evolutionary history of archosaurian muscle activation during locomotion. EMG
20	electrodes were implanted into the muscles of turkeys, pheasants, quail, guineafowl, emus (three
21	age classes), tinamous and juvenile Nile crocodiles across 13 different appendicular muscles.
22	Subjects walked and ran at a range of speeds both overground and on treadmills during EMG
23	recordings. Anatomically similar muscles such as the lateral gastrocnemius exhibited similar EMG
24	patterns at similar relative speeds across all birds. In the crocodiles, the EMG signals closely matched
25	previously published data for alligators. The timing of lateral gastrocnemius activation was relatively
26	later within a stride cycle for crocodiles compared to birds. This difference may relate to the
27	coordinated knee extension and ankle plantarflexion timing across the swing-stance transition in
28	Crocodylia, unlike in birds where there is knee flexion and ankle dorsiflexion across swing-stance. No
29	significant effects were found across the species for ontogeny, or between treadmill and overground

<u> </u>	.ff	2
ι	111	2

30	locomotion. Our findings strengthen the inference that some muscle EMG patterns remained
31	conservative throughout Archosauria: for example, digital flexors retained similar stance phase
32	activity and M. pectoralis remained an "anti-gravity" muscle. However, some avian hindlimb muscles
33	evolved divergent activations in tandem with morphofunctional changes such as bipedalism and
34	more crouched postures, especially M. iliotrochantericus caudalis switching from swing to stance
35	phase activity and M. iliofibularis adding a novel stance phase burst of activity.
36	Keywords: morphology, neural control, musculoskeletal system, evolution, locomotion.
37	Research Highlights
38	Crocodilians show appendicular muscle activity patterns linked to ancestral conservatism. Birds
39	show consistent differences from the ancestral state, which may have been inherited from
40	dinosaurian ancestors after the Triassic archosaurian divergence.
41	
42	Introduction

Animals move using coordinated patterns of muscular activity stimulated by motor neurons. The
electrical signals associated with neuromuscular excitation and thence activation can be obtained
using electromyography (EMG). The relative amplitudes and timings of EMG signals can be used to
(qualitatively) approximate muscle force (Roberts and Gabaldón, 2008). Integrating EMG with
kinematic data and anatomical information facilitates interpretation of the individual function of
muscles (e.g. Roberts and Gabaldón, 2008; Carr et al., 2011).

49

50 Collecting EMG from non-human animals is difficult because surface EMG require ideal conditions 51 (thin skin, minimal skin motion, clean attachment sites, conductive gels, good adhesion, etc.) which 52 are not readily achieved in many animals. As such, the majority of EMG data in animals have been 53 collected by surgically implanted electrodes. Probably because of the difficulties inherent in

54 collecting EMG data from appendicular muscles during locomotion and ethical priorities to minimize 55 the number of invasive animal experiments, only a small range of non-mammalian amniote taxa 56 (and muscles) have been studied. These studies include birds (Jacobson and Hollyday, 1982; Gatesy, 57 1994, 1999; Daley and Biewener, 2003, 2011; Marsh et al., 2004; Daley et al., 2009; Ellerby and 58 Marsh, 2010; Gordon et al., 2015), but also alligators (Gatesy, 1991, 1994, 1997; Reilly et al., 2005), 59 caiman (Gatesy, 1994), turtles (Rivera et al., 2011; Rivera and Blob, 2013), and lizards (Jenkins and 60 Goslow, 1983; Reilly, 1995; Higham and Jayne, 2004; Foster and Higham, 2014, 2017). The majority 61 of these works have focussed on using these species as models for specific mechanistic questions. 62 However, a few studies have attempted to understand muscle activation patterns with the explicit 63 goal of reconstructing the evolutionary diversification of limb motor function (Gatesy and Dial, 1993, 64 1996; Gatesy, 1994, 1999; Rivera and Blob, 2013).

65

66 Better understanding of the relationships between morphology and muscle activity will enable 67 prediction of function for animals (whether extant or extinct) for which no data exist. Such 68 predictions are particularly important for lineages characterized by major changes in functional 69 disparity, such as the Archosauria ("ruling reptiles"; Crocodylia, birds/Aves, and all descendants of 70 their most recent common ancestor). For such clades there is a need to better predict function from 71 form. Archosauria is a clade that diversified first during Triassic period ~250 Mya, evolving a wide 72 variety of forms including small- and large-bodied, sprawling/erect-limbed, quadrupedal/bipedal, 73 aquatic/amphibious/terrestrial and flightless/flying. EMG data from extant archosaurs, particularly 74 the timings of activation during the stride cycle, have been used to infer locomotor changes across 75 Archosauria as a whole (Gatesy, 1994, 1999; Gatesy and Dial, 1996; Hutchinson and Gatesy, 2000). 76 The taxonomic sampling of these EMG data remains somewhat sparse, prompting the question of 77 whether variation in EMG patterns within extant Crocodylia or Aves might alter inferences of 78 neuromotor evolution in Archosauria.

80 Available hindlimb EMG data for birds are largely restricted to guineafowl (e.g. Gatesy, 1999; Marsh 81 et al., 2004; Higham et al., 2008; Daley et al., 2009; Ellerby and Marsh, 2010; Carr et al., 2011; Daley 82 and Biewener, 2011; Gordon et al., 2015) and domestic chickens (e.g. Jacobson and Hollyday, 1982; 83 Bradley and Bekoff, 1992), although some data exist for wild turkeys (Roberts and Gabaldón, 2008), 84 mallard ducks (Anas platyrhynchos) (Biewener and Corning, 2001) and pigeons (Gatesy and Dial, 85 1993, 1996). These data have revealed some consistent patterns of muscle activity such as co-86 activation of muscle pairs (e.g. M. flexor cruris lateralis pars pelvica (hip extension and knee flexion) 87 and M. gastrocnemius pars lateralis (knee flexion and ankle extension) (Ellerby and Marsh, 2010). 88 Almost all avian taxa for which hindlimb EMG data currently exist belong to the clade Galliformes 89 (except mallard ducks – Anseriformes; and pigeons – Columbiformes), a useful model system 90 because they are relatively terrestrial and athletic compared to many other species belonging to the 91 speciose avian clade Neognathae. Hence, almost all present understanding of neuromuscular control 92 of hindlimb function in crown-clade Aves is based upon the assumption that Galliformes represents 93 the typical pattern for all or most birds. This assumption seems reasonable, but it deserves further 94 testing with more data from some of the 9000+ species of extant Aves.

95

96 Importantly, there have been no hindlimb EMG studies of the sister group of Neognathae, the 97 Palaeognathae. The palaeognaths include highly specialized, terrestrial, long-limbed (cursorial) forms 98 such as ostriches, emus, rheas, cassowaries and kiwis, but also the tinamous. Tinamous are of 99 particular interest because they are more similar to "ancestral avian" morphology compared to 100 other paleognaths, with small body size and retained flight capability, and are perhaps even more 101 plesiomorphic in locomotor function than many galliforms (Yonezawa et al., 2017). Prior studies 102 have simulated hindlimb muscle activities in larger palaeognaths (emus (Goetz et al., 2008) and 103 ostriches (Rankin et al., 2016)), which would benefit from further data for experimental validation.

Here, we present hindlimb EMG data during walking and running from two palaeognath species
(elegant-crested tinamous - *Eudromia elegans*, and emus - *Dromaius novaehollandiae*), and four
galliform species (helmeted guineafowl - *Numida meleagris*, wild American turkey - *Meleagris gallopavo*, common pheasant – *Phasianus colchicus* and bobwhite quail - *Colinus virginianus*). We
aim to test whether morphologically conserved hindlimb muscles function similarly across a broad
range of Aves indicated by activation during the same phases of the stride cycles at similar speeds.

110

111 Birds span the range of ontogenetic strategies from altricial to precocial, with precocial birds 112 appearing to be miniatures of their adult form. In chickens (precocial), the neural pathways that 113 drive locomotion appear to arise before hatching, enabling them to walk within hours of hatching 114 (Bekoff, 1976; Bekoff et al., 1987; Bradley et al., 2014). There may also be developmental changes in 115 neural control and muscle activity within birds as they grow (e.g. Tobalske et al., 2017), similar to 116 that seen in certain turtle species (Blob et al., 2008). We measured ontogenetic variation of post-117 hatching neuromuscular control within emus from young birds (< 4kg) to adults (>30kg), for 118 comparison with existing data on ontogenetic scaling of limb muscles (Lamas et al., 2014) and 119 ontogenetic changes of EMG patterns in chickens post-hatching (Bekoff, 1976; Bekoff et al., 1987; 120 Bradley et al., 2014).

121

Finally, we broaden our study's perspective to cover extant Archosauria by including novel EMG data
from Nile crocodiles (*Crocodylus niloticus*). Similarly to birds, all knowledge of appendicular
neuromuscular control in Crocodylia is based on the less diverse subclade Alligatoroidea (Gatesy,
1991, 1994, 1997; Reilly et al., 2005); almost exclusively *Alligator mississippiensis*. By adding EMG
measurements from a representative taxon within the more diverse clade Crocodyloidea, we will
test if similar EMG patterns hold for Crocodylia as a whole, or even more broadly within Archosauria,
Sauria or Tetrapoda (*vide* Gatesy, 1994, 1999). Furthermore, work on birds (domestic chickens)

comparing overground and treadmill locomotion show minor differences in EMG patterns (Jacobson
and Hollyday, 1982), so we test whether this is the case in crocodilians too.

131

Overall, we aim to use this extensive dataset on muscle activity patterns to revisit the questions raised by Gatesy (1994, 1999) about how much diversity exists in the neuromuscular control of locomotion among archosaurs. This comparative dataset will have intrinsic value in applications to other archosaurs; both extant and extinct (e.g. Hutchinson and Gatesy, 2000; Rankin et al., 2016).

136

## 137 Methods

All species, numbers of individuals used, ontogenetic stage, sexes and body masses are listed inTable 1.

140 Ethics

EMG data collection with Nile crocodiles and Elegant-crested tinamous, and the guineafowl and pheasant procedures, were conducted at the RVC Structure and Motion Laboratory under two different project licences approved by the college's Ethics and Welfare committee and granted by the Home Office (United Kingdom). Bobwhite quail and wild turkey data were collected at the Concord Field Station of Harvard University, following procedures licensed and approved by the Harvard Institutional Animal Care and Use Committee in accordance with the guidelines of the National Institutes of Health and the regulations of the United States Department of Agriculture.

148 Surgical procedures

149 For all species bipolar EMG electrodes were constructed of two strands of 0.004 inch diameter

150 platinum pure TC grade (100896) insulated by heavy poly-nylon (HPN) (California Fine Wire

151 Company, CA, USA) soldered to a connector. The free ends of the electrodes had a staggered 1mm

exposed wire region spaced 1.5mm apart. The electrodes were implanted under surgical anaesthesia
appropriate for that species (see details below). Surgeries involved: 1) making skin incisions over the
locations of electrode placement, 2) intramuscular implantation of fine-wire bipolar electrodes, 3)
subcutaneous tunnelling of electrodes to a connector on the dorsum or proximal hindlimb, 4)
closure of incisions and 5) peri- and post-operative administration of analgesia. The recorded
muscles (and their in-text abbreviations) are in Tables 2 and 3.

158

159 *Emu* 

160 Six emus were anaesthetised either using mask inhalation of 5% isoflurane for the chicks, or using 161 intramuscular injections of xylazine (3mg/kg) and ketamine (15mg/kg) to the left caudolateral shank 162 muscles for the juveniles and adults. After inductions, the birds were intubated with an endotracheal 163 tube and maintained at an adequate surgical anaesthetic plane with a variable concentration of 164 inhaled isoflurane. Breathing, heart rate and body temperature were monitored throughout surgery. 165 The feathers in the surgical field were clipped and incisions were made for electrode implantation. 166 The EMG electrodes were successfully implanted into M. Iliotrochantericus caudalis (ITC), M. 167 iliotibialis lateralis pars postacetabularis (ILPO), M. iliofibularis (ILFB) and M. gastrocnemialis pars 168 lateralis (GL) (Figure 1). All wires exited via a skin incision caudal to the femoral trochanteric crest of 169 the right pelvic limb. After surgery, animals were rested in their habitual pen and administered non-170 steroidal anti-inflammatories (meloxicam 1.5mg/kg, three times a day) until data collection was 171 completed. Birds were assessed for discomfort before and throughout data collection; which started 172 24 hours post-surgery; studies were postponed or interrupted if the animals appeared distressed or 173 lame.

174

175 Other birds

176 The guineafowl, pheasant, quail and turkey all underwent surgical procedures that have been 177 described previously (Daley and Biewener, 2003, 2011; Daley et al., 2009), with the birds 178 anaesthetised using isoflurane delivered via a mask. The tinamous followed a similar method (see 179 supplementary information for a more detailed protocol), although general anaesthesia was induced 180 using intramuscular injection of 0.075 mg/kg Ketamine (Ketamidor, Chanelle UK) and 22mg/kg 181 medetomidine (Sedastart, Animalcare UK) into the right pectoral muscle, and maintained using inhaled sevoflurane using a non-cuffed endotracheal tube throughout the remainder of the 182 183 procedure. The surgical field was plucked of feathers and sterilised, and incisions were made over 184 the target muscles. The EMG electrodes were then implanted into the target muscles, while 185 connected to a micro-connector placed on the bird's back. The electrode leads were passed 186 subcutaneously from a 1-2cm incision over the synsacrum to the larger primary incision (4-5cm) over 187 the right lateral shank. Bipolar electrodes were constructed of 0.1mm diameter silver fine-wire 188 (California Fine Wire, Inc., Grover Beach, USA) with 0.5-1.0mm bared tips, and 5-8mm spacing. 189 Electrodes were emplaced using a 23 gauge hypodermic needle, and secured to the muscle using 5-0 190 silk suture; then skin incisions were closed using 3-0 silk. The birds were given analgesia every 12 191 hours and antibiotics every 24 hours. Experimental recordings took place over the next 1–3 days for 192 most birds, but the tinamous were given six days to recover due to their potential sensitivity to 193 stress (pers. obs.,) perhaps due to their relatively small hearts (Altimiras et al., 2017).

194

For all birds, the M. gastrocnemius pars lateralis (GL) was successfully implanted. Additionally, the M. flexor perforatus digiti IV (DFIV) was implanted in both the guineafowl and turkey. Some birds also had implantations into uniquely measured muscles: in the turkey M. flexor cruris lateralis pars pelvica (FCLP), in the guineafowl M. femorotibialis lateralis (FMTL) and in tinamous M. fibularis longus (FL). The basic anatomical positions of all of these muscles are shown in Figure 1, and their approximate actions listed in Table 2.

Cuff	9
------	---

2	n	1
Z	υ	т

#### 202 Nile Crocodiles

203 The anaesthetic procedure is covered in detail in Monticelli et al., (2019), but is briefly outlined here. 204 General anaesthesia was induced using a combination of medetomidine (Sedastart, Animalcare Ltd., 205 York, UK; 0.2 mg kg<sup>-1</sup>) and ketamine (Ketamidor, Chanelle, Berkshire, UK; 10 mg kg<sup>-1</sup>) intramuscularly 206 in the left triceps brachii muscle. After anaesthetic induction, the crocodiles were intubated using an 207 uncuffed endotracheal tube and anaesthesia was maintained using sevoflurane (SevoFlo, Zoetis, 208 Belgium) in oxygen. Intramuscular meloxicam (Metacam, Boehringer Ingelheim, DE; 0.2 mg/kg) was 209 administered in the perioperative period. Active warming was provided by either HotDog® 210 (Augustine Surgical, Eden Prairie, MN, USA) or Bair Hugger® (3M, Maplewood, MN, USA) systems. 211 212 Five incisions, 1-2cm long, were made over the right ilium, proximolateral aspect of the tail, cranial 213 thigh, and caudal and cranial aspects of the lateral shank to enable visualisation and intramuscular 214 implantation for the four hindlimb implants. A further six incisions were made to access four 215 forelimb muscles, with incisions at the scapula, cranial and caudal aspects of the upper arm, medial 216 aspect of the lower arm, lateral aspect of the thorax, and ventral aspect of the thorax. Through these 217 incisions, the muscles were implanted. After post-mortem, the muscles from which data were 218 collected were confirmed to be M. transversus perinei (TP), M. iliotibialis 2 (IT2), M. gastrocnemius 219 externus (GE), M. flexor digitorum longus (FDL) of the hindlimb and M. pectoralis (PEC) of the 220 forelimb (Figure 1, Table 3).

221

The EMG electrode connector was anchored by suturing to two scutes near the dorsal-most incision (iliac or scapular). Each pair of electrode wires was then subcutaneously tunnelled to their respective insertion sites. Tunnelling was achieved subcutaneously using a section of size 3 (internal diameter) 225 uncuffed PVC endotracheal tubing and a looped guide wire. The electrodes were implanted using 226 the sew-through method and secured with two simple-interrupted sutures using 3-0 vicryl to 227 prevent both translation and rotation of the wires post-surgery. The excess wiring was pulled back 228 through to the dorsal incisions where it was coiled and tucked back into the incision site. Each 229 incision was then flushed with lidocaine and then closed using everted mattress stitches to prevent 230 wound contamination in the water within the enclosures. The anaesthesia was discontinued and 231 atipamezole (1 mg kg<sup>-1</sup>) (Sedastop, Animalcare, UK) was administered intramuscularly in the left M. 232 triceps brachii, and repeated after 30 minutes in case of residual sedation. The crocodiles were then 233 given at least two days to recover in their enclosures before any data were collected

234

#### 235 Experimental protocol

236 The tinamous were placed on a Starkerhund treadmill (Terraglione di Vigodarzere, Italy) within a box 237 with transparent acrylic sides to allow visualisation of the footfalls, and which had an opening for the 238 EMG wires. Trials ranged from 30s to 60s, with at least a 60s break between trials. The treadmill speed varied from 0.1ms<sup>-1</sup> to 0.45ms<sup>-1</sup>; faster speeds were not safely achievable with the birds. The 239 240 trials were initiated using a trigger system that created a short light flash that could be seen by the two Hero 3+ GoPro cameras (San Mateo, CA, USA) recording at 60Hz which were used to capture the 241 footfall patterns of the animals during locomotion. Trials were maximally 60s long, although usually 242 243 far shorter, with at least 60s recovery between trials. The birds were in the experimental area for a 244 maximum of 1hr before being returned to their enclosure. A total of 64 trials were collected for the 245 two individuals, with the resulting data summarised in Table 5.

246

248 Emu experimental trials were conducted overground in a corridor of ~90cm width enclosed by wire 249 netting for the younger individuals, and metal fencing for the adults. Due to the wired EMG 250 implants, cable length limited the maximum length of the runway. Cable length was ~5m for the 251 youngest birds and 9m for the two older groups. All wires were tethered along a sliding pulley 252 system (suspended >1m off the ground) which kept the implant cables from dragging on the floor 253 and interfering with gait. The floor of the runway was instrumented with eight Kistler forceplates 254 (0.6x0.9m; model 9287B, Hook, Hampshire, UK), which were used to obtain timings of footfalls. The 255 emus were also marked with polystyrene hemispheres covered with infrared-reflective tape 256 (Scotchlite 8850; 3M, Manchester, UK) (1cm diameter for the youngest, 2cm diameter for the older 257 individuals) for joint motion analysis for another study (Lamas, 2015), which included two dorsal 258 midline body markers used here for obtaining locomotor velocities via a Qualisys Oqus 500 six-259 camera system recording at 250Hz (Qualisys AB, Göteborg, Sweden). Across the six individuals, 405 260 trials were completed, and the resulting trials are listed in Table 6.

261

The turkey (2 individuals, 5 trials), quail (2 individuals, 6 trials), guineafowl (2 individuals, 6 trials), and pheasant (1 individual, 1 trial) all ran on a custom-built treadmill, with a slatted black rubbercoated steel belt with a 55.8×172.7 cm running surface. The treadmill speeds were selected to achieve an approximately similar dimensionless speed (see below) of 1.25 across species (Table 5). Dimensionless speed is the square root of the Froude number (Alexander and Jayes, 1983):

267 
$$u = \frac{v}{\sqrt{(g \cdot l)}}$$

where *u* is dimensionless speed, *v* is velocity (ms<sup>-1</sup>), *g* is acceleration due to gravity (9.81ms<sup>-2</sup>) and *l* is standing (or mid-stance) hip height (in metres). Dimensionless speed usually assumes geometric similarity (Alexander and Jayes, 1983); however, dimensionless speed holds reasonably well across animals that use similar locomotor modes even if not strictly geometrically similar (see Daley and Birn-Jeffery, 2018) and references therein for a thorough review). The turkey, quail and quineafowl
were recorded using a Photron camera (Photron Europe Ltd., West Wycombe, UK) at 125Hz, whilst
the pheasant was recorded using Qualisys cameras (as per the emus above) at 125Hz.

275

276 The crocodiles were captured from their enclosures and their mouths were taped to prevent injury 277 to themselves or handlers, or damaging their EMG wires. The crocodiles were then either placed on 278 a Starkerhund treadmill (within an acrylic-sided enclosure to prevent the animals escaping), or on a 279 custom-made wooden runway (0.38x0.40x2.44m). Both the treadmill enclosure and custom wooden 280 runway had openings in the roof to allow the wires to exit to be connected to the EMG amplifiers. 281 The crocodiles were motivated to walk by stimulating the tail with a broom as needed. The hardware 282 otherwise was the same as that used with the tinamou. Trials were maximally 60s long, although 283 usually far shorter, with at least 60s recovery between trials. Across four individuals, a total of 160 284 trials were collected, with details of collected data in Table 4.

285

#### 286 EMG recordings

Each of the sockets on the animals was connected via lightweight shielded cables to GRASS preamplifiers (P511, Natus Neurology Inc., Pleasanton, CA, USA). EMG signals remained at a constant
amplification throughout data collection with a low-pass (10Hz for most birds; 30Hz for emus,
tinamous and crocodiles) and a high-pass (10Hz) filter. The EMG signals were sampled at 2500Hz
(emu) or 5000Hz (all other species). Signals were amplified between 1000 and 10000 times, but this
varied between and within data collection sessions and individuals as required to obtain visible
signals.

294

295 Data processing

Footfall events (foot on and off times) were manually recorded from the videos for the crocodiles,
tinamous, guineafowl, quail, turkey and pheasant for each trial. The emu footfall timing pattern was
determined by analysing the forceplate data, with foot on and off timings linked to the force traces
(recorded at 1000Hz; automatically filtered using a low-pass filter at 100Hz; threshold for foot on/off
events = 1 % body weight). Custom scripts in Matlab software (MathWorks, Natick, MA, USA) were
used for all post-processing.

302

303 The tinamous and crocodiles on the treadmill moved at three different speeds, the lowest being 304 driven by an electric drill turning the drive wheel, the other two (approximately 0.5, 0.7mph) driven 305 by the treadmill motor. The belt speed was calibrated from a video based on the movement of a 306 mark on the treadmill belt relative to a point at a known distance on the treadmill frame. For the 307 crocodiles on the runway, locomotor speed was measured by tracking the shoulder scutes (which 308 had the least lateral movement relative to direction of movement) across 20cm, using a dorsally 309 placed camera atop the runway. Emu speeds were calculated by tracking the cranial dorsal body 310 marker in 3D space, then extracting the horizontal displacements and dividing by the time elapsed.

311

Data were then cut into individual steps based on footfall timings extracted from video or forceplate data from each species, described above. Each EMG sequence was filtered and then rectified. The data were bandpass-filtered with a low-pass cutoff between 50-90 Hz and a 12-order Butterworth filter applied, although the emu data underwent a 0.5 Hz low-pass filter to remove an underlying noise waveform before being filtered as the other data (see Figure 2 for representative filtered EMG signals).

319 In species with multiple speeds, dimensionless speed was used for comparisons. As the emu trials 320 spanned the largest range of dimensionless speeds, they were grouped into 0.2*u* bins from 0.3-1.5*u*. 321 These bins covered all of the ontogenetic stages of the emu and overlapped with the recorded speed 322 range of other studied bird species except for the tinamou, which never reached a dimensionless 323 speed greater than 0.3. The crocodiles' speeds were also normalised to dimensionless speed where I 324 is total hindlimb length instead of hip height due to the variety of postures (from sprawling to 325 upright) that they used. For each species, the rectified sequences for each grouping (Table 2,3) were 326 scaled to the same length with "foot-on" being 0 and 100 (equivalent to a full stride). The average 327 and 95% confidence intervals for each of these groupings were then calculated using custom Matlab 328 code. EMG onset was deemed to be where there were peaks beyond 10% of the baseline.

329

For one muscle (M. gastrocnemius lateralis/externus; GL/GE) there were sufficient data for statistical tests of gross similarity of EMG signals across Aves and Crocodylia. An inter-species cross-correlation analysis was carried out upon the average rectified EMG sequences for comparable speeds of 1.1-1.3*u* (or maximum speeds for tinamous and crocodiles), using custom Matlab scripts. Due to the small sample sizes in terms of both numbers of individuals and number of trials, no additional statistics were carried out.

336

337 Results

338 Birds

339 <u>Emus</u>

341

340 There were no major differences between the age groups in terms of muscle activation timings

within the stride cycles. However, the baby emus may have had a slightly broader range of

- 342 activations for each muscle group relative to the older individuals. With so few individuals, it is
- 343 difficult to resolve whether this difference related to individual variance or age (Figure 3).

344

345 *ITC* (Figure 3, S1)

346 The M. iliotrochantericus caudalis was active from late swing through to mid-stance . However, at

slower relative speeds (below 1.1*u*), the late swing and early stance activations were at lower levels,

348 and the peak activation occurred during mid-stance. At the fastest speeds, the activation peaked in

349 early stance.

350

351 *ILPO* (Figure 3, S2)

352 The M. iliotibialis lateralis pars postacetabularis had variable activation with speed. At slower speeds

353 (<0.7 *u*), the EMG signals occurred at a fairly consistent level of activity from the end of swing

through late-middle stance phase. At faster relative speeds (0.7-0.9 *u*), the EMG signal peaked

during mid-stance, with lower activity around late swing and later stance. At the fastest speeds (0.9-

1.1 and 1.1-1.3 *u*), there was a discontinuity between the signal in late swing and the large peak at

357 mid-stance; consistent with two bursts of activity.

358

359 *ILFB* (Figure 3, S3)

The M. iliofibularis displayed peak activation during early stance, but with activity persisting fromlate swing to mid-stance.

362

363 GL (Figure 3, S4)

Cuff 1
--------

364	The M. gastrocnemius pars lateralis of emus, like the ITC and ILFB, was active from late swing
365	through mid-stance phase at most speeds. In the youngest emus, EMG activity extended through
366	stance and into early swing at $u$ from 0.3-0.5. At higher values of $u$ , the initial muscle activations for
367	all ages became increasingly earlier, so activation occurred more consistently during late swing and
368	ended earlier in stance, with higher relative activations and a smaller range as a proportion of total
369	stride time.
370	
371	<u>Tinamous</u> (Figure 4, S5)
372	GL
373	Overall, the M. gastrocnemius pars lateralis EMG signals were similar across the small range in
374	speeds, with activity beginning in late swing and continuing through early stance phase, with a
375	reduced mid-late stance signal at faster speeds.
376	
377	FL
378	From the one trial of 7 strides at $0.1 \text{ms}^{-1}$ (0.06 <i>u</i> - a very slow walk), the M. fibularis longus showed
379	low level EMG activity from foot on through to late-middle stance phase.
380	
381	Galliform birds
382	FCLP
383	The M. flexor cruris lateralis pars pelvica was only measured in turkeys, and was active through
384	stance phase, with a peak in mid-stance (Figure 5, S6).
385	

	Cuff 17
386	FMTL
387	The M. femorotibialis lateralis was only measured in the guineafowl, and was active from late swing
388	through to late stance phase (Figure 6, S7).
389	
390	GL
391	Across the quail, pheasants, guineafowl and turkey, the lateral gastrocnemius showed a similar
392	overall pattern of activity, with the primary burst of muscle activity occurring from late swing to
393	early mid-stance phase, with peak activity early in stance (Figure 7A-D, S8).
394	
395	DFIV
396	The M. flexor perforatus digiti IV was measured in turkeys and guineafowl, and showed similar
397	activity from late swing through early stance phase; as in the GL (Figures 5-6, S6-S7).
398	
399	
400	Crocodiles (Figure 8, S9)
401	PEC
402	The M. pectoralis for the Nile crocodiles showed low-level activation through mid-stance phase, with
403	maximal activation in late stance. Unlike the TP (below), the pattern was not shifted earlier in the
404	cycle at faster speeds; instead, there was relatively shorter period of activation. For relatively similar
405	speeds, there was no apparent difference between EMG signals for animals on treadmill or runways.
406	

407 TP

408	The M. transversus perinei was active through early to mid-stance phase, with peak activity from 20-
409	50% of the stride cycle. At faster speeds, the TP became active earlier, including late swing phase
410	(Figure 8).
411	
412	IT2
413	The M. iliotibialis 2 was active throughout most of the stance phase, with the greatest signals around
414	30% of total stride cycle.
415	
416	GE
417	At 0.35-0.45ms <sup>-1</sup> walking speeds, the M. gastrocnemius externus was active during mid-late stance,
418	becoming active into early swing phase at faster speeds.
419	
420	Our cross-correlation analysis of mean EMG timings for the GE of Crocodylia and GL of Aves showed
421	that these were most similar for Aves, and distinct for Crocodylia (Table 7). Interestingly, the
422	maximal correlations of timings for Crocodylia were more similar to those of Palaeognathae and
423	quails, whereas guineafowl and turkey were most similar to each other; with pheasant values in
424	between these. However, these similarities in maximal correlations were not so evident in the
425	offsets of EMG timings, which were ~18-27% of a stride earlier in swing (start-signal) and stance
426	(end-signal) for all Aves vs. Crocodylia.
427	
428	

429 Discussion

We have presented a compilation of the largest dataset of electromyographic data for archosaurs to
date, including the first for emus and tinamous, thus adding palaeognathous birds to the existing
literature for birds and Nile crocodiles to the published data for Crocodylia. Below, we consider our
avian data first, then crocodilian EMG data, then all data in the broader context of archosaurian
neuromuscular evolution.

435

Published hindlimb EMG data from birds to date are primarily from guineafowl (*Numida*) and
domestic chickens (Jacobson and Hollyday, 1982; Gatesy, 1999; Ellerby and Marsh, 2010), as well as
wild turkeys (Roberts and Gabaldón, 2008), although some important data also exist for pigeons
(Gatesy and Dial, 1993, 1996). The guineafowl data presented here compare well with previously
published guineafowl data, both in terms of patterns and timings for the GL and DFIV muscles;
bolstering confidence in our results (Gatesy, 1999; Daley and Biewener, 2003; Gordon et al., 2015).

442

443 Prior to this study, there were no palaeognath EMG data, and the only relevant data were derived 444 from a musculoskeletal simulation studies of emus (Goetz et al., 2008) and ostriches (Rankin et al., 445 2016), which predicted muscle activations for walking and running. Goetz et al. (2008) only depicted 446 muscle forces rather than activations, and solely for the stance phase—but for 40 muscles or 447 subdivisions thereof. Rankin et al. (2016) reported activation timings (e.g. see their figure 3 for 16 448 muscles) that generally match the additional palaeognath EMG data we present here, with the 449 exception of the ITC(p) which consistently had a secondary activation during swing phase in the 450 simulations. Whilst some small peaks were evident in our ITC data for the adult emus around foot-451 off at slower speeds, these are likely a result of noise, and no peaks indicative of secondary 452 activations in mid-swing were found for running emus (u >1.0) (Figure 3, S1). This discrepancy 453 between the simulated muscle activations and EMG data was also found when compared to 454 guineafowl data (Daley et al., 2009). Furthermore, the simulated muscle activations may be

misleading because these rely on Hill-type muscle models that presume basic relationships between
neural excitation, muscle activation and force, whereas in reality these relationships may be more
complex (e.g. Askew and Marsh, 1998; Perreault et al., 2003; Millard et al., 2013).

458

459 Did we observe ontogenetic variation within our emu data? Whilst variation in EMG signals through 460 ontogeny has been found in ovo (e.g. Watson and Bekoff, 1990), very little research has been done 461 on post-hatching birds (see Jacobson and Hollyday, 1982; Muir et al., 1996) as neural controls 462 appear to establish early within embryos. However, in chukar partridges (Alectoris chukar) there is variation between young and adult birds in EMG patterns for pectoralis muscle EMG signals during 463 464 wing-assisted incline running, with younger birds having longer periods of activation of their muscles 465 relative to the adults (Tobalske et al., 2017). We found a similar pattern of longer activation times in 466 younger emus' leg muscles, although the variation between the different ages was small. These 467 similarities are in contrast to turtles. Juvenile turtles can undergo significant changes in muscle 468 activity patterns through ontogeny, usually decreasing the number of activations for muscles during 469 the stride cycle from two to one but sometimes profoundly changing the timing of muscle activation. 470 For example, across the ontogeny of juveniles to adults the femorotibialis muscle's activation 471 transitions from recovery (swing) to thrust (stance) phase in swimming (Blob et al., 2008).

472

The M. gastrocnemius pars lateralis (GL) has the EMG patterns that are most widely studied across avian species, and serves as a useful as a reference muscle because it retains similar anatomy (origin on the lateral distal femoral condyle and insertion on the tarsometatarsus) and locomotor function (primarily ankle extension but also knee flexion) in terrestrial gait across species. Here we found that the GL's EMG activity patterns of the palaeognathous birds were almost exactly the same as those for the neognaths during walk/run behaviours (Figure 7, Table 7). Activity began in late swing phase and continued throughout much of stance, but at faster speeds became increasingly concentrated in 480 early stance phase, in all birds studied to date. We suggest that across cursorial, ground birds as a 481 whole, muscle activation patterns are likely to be conserved for morphologically and functionally 482 similar muscles. More broadly, even GL activations during walking in the more aquatically specialized 483 mallard duck (Anas platyrhynchos) are similar to the species reported here (Biewener and Corning, 484 2001). Thus GL muscle activation patterns appear to be generally conserved across Aves and 485 correspond to the expected functional demands inferred from anatomical origins, insertions, joint 486 mobility and moment arms. However, there may be differences in EMG timing of the GL during the 487 stance phase that correlate with differences in limb posture (e.g. Gatesy and Biewener, 1991; Daley 488 and Biewener, 2011; Gordon et al., 2015; Daley and Birn-Jeffery, 2018) and anatomical variations; or 489 perhaps even across regions of the GL.

490

491 As in the previous study of chickens (Jacobson and Hollyday, 1982), regardless of whether the 492 crocodiles were moving within a runway or on a treadmill, their muscle activation patterns were 493 very similar, although the range of speeds assessed was very low, approximately at their natural 494 walking speed (0.1ms<sup>-1</sup>). Previously published data for Crocodylia derive entirely from the American 495 alligator (Alligator mississippiensis) and spectacled caiman (Caiman crocodilus) hindlimb (Gatesy, 496 1994, 1997; Reilly and Blob, 2003); in both cases for the Alligatoroidea lineage. Here, we provide 497 comparable data for the Crocodyloidea lineage (and the first forelimb PEC muscle EMG for 498 Crocodylia). The M. iliotibialis 2 (IT2) of Nile crocodiles had a relatively larger EMG signal than seen 499 in alligators but it occurred with the same timing around mid-stance (Gatesy, 1997). Whilst no 500 filtered signals are published for the M. gastrocnemius externus (GE), summarized timings (Reilly et 501 al., 2005) match very well with the signals seen in the Nile crocodile data at 0.345ms<sup>-1</sup> presented 502 here (Table 7), despite being from different duty factors (0.6 here vs. 0.7 in Reilly et al., (2005)).

504 The M. transversus perinei (TP) overlies the M. caudofemoralis longus (CFL) in Crocodylia (Romer, 505 1923; Frey, 1982; Cong, 1998) and has similar EMG signals in terms of patterns and timings (Gatesy, 506 1997). Its activity has not been measured before, to our knowledge, in Reptilia. An explanation for 507 this similar activity may be that the TP was contracting simultaneously with the CFL. The two 508 muscles also have muscle fibres that run perpendicular to each other (TP dorsoventrally, CFL 509 craniocaudally). Perhaps the TP performs some function of locomotor relevance, limiting bulging of 510 the distal CFL belly near where it narrows into its insertion, or acts similarly to the M. caudofemoralis 511 brevis, which seems to help to change the moment arm of the CFL in lizards (Herrel et al., 2008). 512 However, the TP is very thin and whilst not visibly implanted within the CFL, the electrodes may have 513 been picking up "cross-talk" signals from this much larger muscle. Regardless, these novel data from Nile crocodiles have intrinsic value for understanding function of the CFL and dynamics of the tail-to-514 515 thigh region; and our EMG data are unlike those of the homologous M. caudofemoralis (pars 516 caudalis) of birds which is only (variably) active during late stance at fast speeds (Gatesy and Dial, 517 1993; Gatesy, 1999).

518

519 There are no other published forelimb EMG data for Crocodylia; however, some data exist for turtles 520 (Rivera and Blob, 2010), which may form the sister clade to archosaurs (Hedges and Poling, 1999; 521 Field et al., 2014 but see Gauthier et al., 1988; Lyson et al., 2012), and the perhaps more distantly 522 related Savannah monitor lizard (Varanus exanthematicus) (Jenkins and Goslow, 1983). The M. 523 pectoralis data from the literature are variable, with turtles showing activity from early to late stance 524 phase (Rivera and Blob, 2010, 2013). In contrast, in the monitor lizard, cranial portions of the PEC 525 were active predominantly in swing phase, whilst the middle of the PEC was active at low levels 526 during stance phase. In our Nile crocodile subject, the electrode was inserted into the cranial (i.e. 527 major sternal) portion of the PEC, but had activity through mid-to-late stance. The differences 528 between the species might relate to differences in the role of the pectoralis in resisting

glenohumeral abduction imposed by ground reaction forces, whether due to the "high-walking"postures in crocodiles or the added mass of the shell in turtles.

531

532 What, then, do our EMG data indicate about the evolution of muscle activity in the clade 533 Archosauria? There are scarce overlapping and ideally comparable data, major differences in 534 locomotor biomechanics and some issues with muscle homology (Rowe, 1986; Gatesy, 1994) that 535 are cause for caution. On the other hand, there are clearly corresponding patterns of muscle activity 536 that also match similarities in limb dynamics and/or muscle anatomy for Archosauria (and even other tetrapods) (Figure 9). For example, the large, ventral M. pectoralis would be expected to be an 537 538 antigravity muscle from the anatomy and indeed both Crocodylia (represented by our new data for 539 Crocodylus niloticus) and Aves activate M. pectoralis during their major antigravity functions (i.e. 540 stance phase for the former; downstroke of flight for the latter) (Gatesy and Dial, 1993, 1996; 541 Goslow et al., 2000) as expected for this large, ventrally disposed muscle. This antigravity activity is 542 broadly shared with turtles and Squamata (Varanus exanthematicus: caudal PEC region), consistent with some "neuromotor conservation" at least across the broader amniote clade Sauria (Jenkins and 543 544 Goslow, 1983; Schoenfuss et al., 2010; Rivera et al., 2011; Rivera and Blob, 2013). This apparent 545 conserved activity would support the inference that quadrupedal archosaurs used their PEC muscles 546 to support themselves during locomotion, much as their M. adductor femoris muscles countered 547 abduction of the hindlimbs (Hutchinson and Gatesy, 2000). However, the swing phase activity of the 548 cranial region of PEC in Varanus is cause for caution, as it does not follow the expectation of a 549 ventrally located muscle acting simply as an anti-gravity adductor, and does not match the activity of 550 the comparable region in *Crocodylus*.

551

Our hindlimb EMG data for Crocodylia and Aves indicate broadly similar stance phase activity for
 GE/GL in both groups of extant archosaurs; although unsurprising, this is nonetheless consistent

554 with a conserved antigravity function that would be expected possibly throughout Tetrapoda (e.g. 555 turtles (Schoenfuss et al., 2010), lizards (Reilly 1995, 1998), humans (Dietz et al., 1979), felids 556 (Rasmussen et al., 1978), and salamanders (FPC = medial gastrocnemius)(Ashley-Ross, 1995)). This 557 apparent conservatism also seems to apply to the digital flexors (FDL in the crocodile and DFIV in the 558 guineafowl; the latter homologous to a part of the former (Hutchinson, 2002)), which are also ankle 559 extensors (plantarflexors) with antigravity functions and thus show timings similar to those seen in 560 the GE/GL. Similarly, the IT2/ILPO (here represented by new data for Crocodylus and Dromaius) are 561 homologous muscles for Archosauria (Hutchinson, 2001) and exhibit stance phase activity (earlier in 562 stance/late in swing) in this study and related literature cited above, although our emus had an extra 563 potential burst that may be apomorphic. These data are most parsimoniously interpreted as 564 homologous muscle activity that was ancestral for Archosauria.

565

566 There was one potentially interesting difference in GE/GL timing we observed for Crocodylia vs. 567 Aves: the homologous GE/GL muscles are active only in stance phase in Crocodylia, whereas activity 568 starts in late swing phase in Aves (Table 7, Figure 7). Considering the grossly similar anatomy of 569 these muscles in extant archosaurs, published differences in knee and ankle joint kinematics suggest 570 one possible explanation: that the earlier onset of EMG activity in the avian GL is related to 571 maintaining synchronized knee flexion and ankle dorsiflexion across the swing-stance transition (e.g. 572 (Higham and Nelson, 2008)), unlike in Crocodylia where knee extension and ankle plantarflexion 573 occur from late swing to early stance phase (Gatesy, 1991, 1997; Reilly et al., 2005). This is simply 574 one reasonable speculation that deserves testing against alternatives such as plantigrade foot 575 posture in future studies, but we are unaware of it being previously proffered as an explanation.

576

577 Other EMG data for archosaurian hindlimb muscles are not feasible to compare directly within our
578 dataset. The CFL muscle surely maintained stance phase activity in early archosaurs (Gatesy, 1990).

579 Here, the overlying TP muscle of Crocodylus was sampled rather than the underlying CFL muscle, but 580 the TP has barely been studied in the clade Sauria and deserves further analysis in the context of 581 limb function. The ITC muscle of Aves is homologous with M. iliofemoralis of Crocodylia (Rowe, 582 1986). Our palaeognath EMG data strengthen the hypothesis that there was a switch from swing to 583 stance phase activity of the ITC within the clade Dinosauromorpha, perhaps concurrent with the 584 origin of bipedalism and increased need for hip abductor-based support (rather than adductors) 585 during stance phase (Hutchinson and Gatesy, 2000). Likewise, the late swing and stance phase 586 activity of the ILFB for emus and other birds, vs. mainly swing phase activity in Alligator (Gatesy, 587 1997; Reilly et al., 2005) and Varanus (Jayne et al., 1990), support the conclusion that this muscle 588 added a prominent stance phase burst at some point after the divergence of the 589 dinosauromorph/avian lineage from Archosauria, albeit apparently maintaining a swing phase burst 590 in most birds (Gatesy, 1999). Stance phase activity of the FL in our one tinamou subject and trial, 591 with other avian data (Gatesy, 1994, 1999), support the inference that this activity is ancestral for 592 Aves. Then, considering similar activity of the FL in the lizard Sceloporus (Reilly, 1995), perhaps the FL 593 has conserved stance phase (and perhaps late swing phase) activity across at least Sauria (like the 594 GE/GL) (Reilly, 1998); although EMG data for the FL are lacking in Crocodylia.

595

596 While our data, and synthesis of data from the literature, suggest "conservatism" in muscle 597 activation patterns across Archosauria, Sauria or even more broadly, an explanation for such 598 patterns in terrestrial locomotion remains lacking. The vertebrate sensorimotor control system is 599 plastic and adaptable to allow versatile function; yet retains similar elements of spinal central 600 pattern generators and segmentally arranged reflexes across vertebrates (Grillner and Wallén, 601 1985). Hardwired neural circuitry does not require conserved activation patterns if that circuitry 602 itself enables adaptive feedback (e.g. central pattern generators are well known to be 603 entrained/regulated by sensory feedback and descending control). However, our data here, and

from other recent studies; e.g. turtles (Rivera et al., 2011; Rivera and Blob, 2013); suggest that many patterns of activation broadly are conserved. One potential reason is that the shared functional demands of terrestrial locomotion across all animals (e.g. supporting body weight), combined with shared substrates of actuation and control (muscle and nervous system intrinsic physical properties), mean that most terrestrial animals use similar mechanisms for economic movement, which is reflected in conserved function in muscles that have grossly similar morphology. However, other perspectives have cited reasons to be wary (Smith, 1994; Alfaro and Herrel, 2001).

611

Yet regardless of the cause(s) of a lack of change of motor patterns, such homology is valuable to 612 613 evolutionary biomechanists. Here, we have added to other perspectives on the evolution of 614 appendicular muscle control in archosaurs (Gatesy, 1990, 1999; Gatesy and Dial, 1996; Hutchinson 615 and Gatesy, 2000; Reilly et al., 2005) by showing how a forelimb muscle (PEC) and several hindlimb 616 muscles (IT2/ILPO, ILFB, GE/GL, FDL/DFIV, potentially FL) have maintained similar motor patterns in 617 extant Archosauria, although the avian GL has modified its timing (Figure 9). These similarities were 618 somewhat expected from relatively conservative muscle morphology, and supported prior studies 619 (Gatesy, 1994, 1999), yet the GL timing difference might be less expected. This overall similarity of 620 muscle activations bolsters their usage in validating computer simulations, or otherwise inferring 621 locomotor function, for taxa without available EMG data, whether they are extant archosaurs (Goetz 622 et al., 2008; Rankin et al., 2016) or extinct, as long as the muscles are known or can be inferred 623 (Witmer, 1995). However, the evidence for changes of (or additions to) the motor patterns of avian 624 muscles such as the morphologically transformed ITC and more morphologically conservative ILFB is 625 cause for caution (neuromotor conservation demands to be tested, not assumed (Gatesy, 1994, 626 1999; Smith, 1994; Goslow et al., 2000; Rivera et al., 2011; Rivera and Blob, 2013)), and cause for 627 assembling datasets from more varied taxa and behaviours.

#### 629 Contributions

JRH and MAD conceived the study. ARC, VRA, KBM, LPL and MAD all planned and performed
surgeries on the animals, assisted by CA, PM and LP. ARC, VRA, KBM, LPL, MAD and JRH all carried
out experiments. ARC conducted the data analysis assisted by MAD. ARC wrote the manuscript aided
by JRH and MAD. All authors contributed to reviewing the manuscript and approved the final draft.

634

## 635 Acknowledgements

636 Enrico Eberhard, Peter Bishop and Jorn Cheney all provided advice on Matlab code. We thank La 637 Ferme aux Crocodiles (Pierrelatte, France) for provision of the Nile crocodile subjects. We appreciate the support of the Biological Services Unit at RVC for animal care. We thank Russell Main, Emily 638 639 Sparkes, Sandra Shefelbine and Heather Paxton for help with the experimental data collection for 640 emus, and Jeffery Rankin and James Usherwood for input on that study. Thanks to Alison Tarbell and 641 Sheridan Golding for assistance in quail, turkey and pheasant data collection. This study was supported by funding from The Royal Veterinary College and the European Research Council (ERC) 642 643 under the European Union's Horizon 2020 research and innovation programme (grant agreement 644 #695517). Finally thanks to the anonymous reviewers whose comments greatly improved the manuscript. 645

646

## 647 Conflict of Interests

648 None are declared.

## 649 **References**

Alexander RM, Jayes AS. 1983. A dynamic similarity hypothesis for the gaits of quadrupedal
 mammals. J Zool 201:135–152.

652 653	Alfaro ME, Herrel A. 2001. Introduction: major issues of feeding motor control in vertebrates. Am Zool 41:1243–1247.
654 655 656	Allen V, Molnar J, Parker W, Pollard A, Nolan G, Hutchinson JR. 2014. Comparative architectural properties of limb muscles in Crocodylidae and Alligatoridae and their relevance to divergent use of asymmetrical gaits in extant Crocodylia. J Anat 225:569–582.
657 658 659	Altimiras J, Lindgren I, Giraldo-Deck LM, Matthei A, Garitano-Zavala Á. 2017. Aerobic performance in tinamous is limited by their small heart. A novel hypothesis in the evolution of avian flight. Sci Rep 7:15964.
660 661 662	Ashley-Ross MA. 1995. Patterns of hind limb motor output during walking in the salamander <i>Dicamptodon tenebrosus,</i> with comparisons to other tetrapods. J Comp Physiol A 177: 273- 285.
663 664 665	Askew GN, Marsh RL. 1998. Optimal shortening velocity (V/Vmax) of skeletal muscle during cyclical contractions: length-force effects and velocity-dependent activation and deactivation. J Exp Biol 201:1527–1540.
666 667	Bekoff A. 1976. Ontogeny of leg motor output in the chick embryo: a neural analysis. Brain Res 106:271–291.
668 669 670	Bekoff A, Nusbaum MP, Sabichi AL, Clifford M. 1987. Neural control of limb coordination. I. Comparison of hatching and walking motor output patterns in normal and deafferented chicks. J Neurosci 7:2320–2330.
671 672 673	Blob RW, Rivera ARV, Westneat MW. 2008. Hindlimb Function in Turtle Locomotion: Limb Movements and Muscular Activation across Taxa, Environment, and Ontogeny. In: Wyneken J, Godfrey MH, Bels V, editors. <i>Biology of Turtles</i> . Boca Raton, FL: CRC Press. p. 139–162.
674 675 676	Bradley NS, Bekoff A. 1992. Development of coordinated movement in chicks: II. Temporal analysis of hindlimb muscle synergies at embryonic day 10 in embryos with spinal gap transections. J Neurobiol 23:420–432.
677 678	Bradley NS, Ryu YU, Yeseta MC. 2014. Spontaneous locomotor activity in late-stage chicken embryos is modified by stretch of leg muscles. J Exp Biol 217:896–907.
679 680	Carr JA, Ellerby DJ, Marsh RL. 2011. Function of a large biarticular hip and knee extensor during walking and running in guinea fowl ( <i>Numida meleagris</i> ). J Exp Biol 214:3405–3413.
681 682	Cong L. 1998. Yangzi e da ti jie pou = The gross anatomy of Alligator sinensis (Fauvel). Di 1 ban. ed. Beijing: Ke xue chu ban she.
683 684 685	Daley MA, Biewener AA. 2003. Muscle force-length dynamics during level versus incline locomotion: a comparison of in vivo performance of two guinea fowl ankle extensors. J Exp Biol 206:2941–2958.
686 687 688	Daley MA, Biewener AA. 2011. Leg muscles that mediate stability: mechanics and control of two distal extensor muscles during obstacle negotiation in the guinea fowl. Philos Trans R Soc Lond B, Biol Sci 366:1580–1591.
689 690	Daley MA, Birn-Jeffery A. 2018. Scaling of avian bipedal locomotion reveals independent effects of body mass and leg posture on gait. J Exp Biol 221.

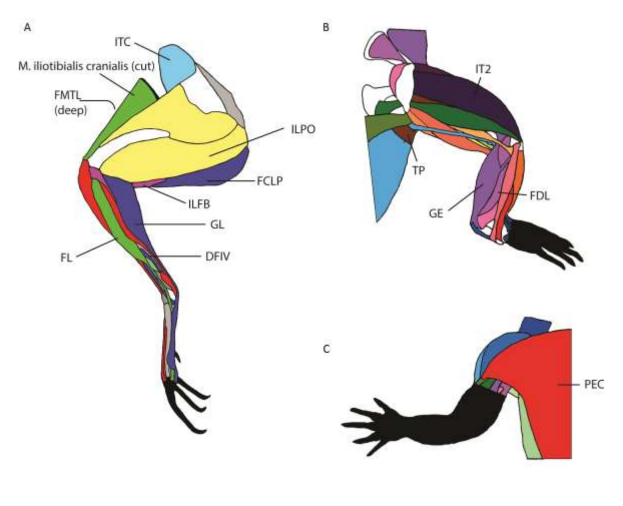
- Daley MA, Voloshina A, Biewener AA. 2009. The role of intrinsic muscle mechanics in the
   neuromuscular control of stable running in the guinea fowl. J Physiol (Lond) 587:2693–2707.
- 693 Dietz V, Schmidtbleicher D, Noth J. 1979. Neuronal mechanisms of human locomotion. J
   694 Neurophysiol 42:1212–1222.
- Ellerby DJ, Marsh RL. 2010. The mechanical function of linked muscles in the guinea fowl hind limb. J
   Exp Biol 213:2201–2208.
- Field DJ, Gauthier JA, King BL, Pisani D, Lyson TR, Peterson KJ. 2014. Toward consilience in reptile
   phylogeny: miRNAs support an archosaur, not lepidosaur, affinity for turtles. Evol Dev
   16:189–196.
- Foster KL, Higham TE. 2014. Context-dependent changes in motor control and kinematics during
   locomotion: modulation and decoupling. Proc Biol Sci 281:20133331.
- Foster KL, Higham TE. 2017. Integrating gastrocnemius force-length properties, in vivo activation and
   operating lengths reveals how *Anolis* deal with ecological challenges. J Exp Biol 220:796–806.
- Frey E. 1982. Ecology, locomotion and tail muscle anatomy of crocodiles. Neues Jahrbuch fur
   Geologie und Paläontologie Abhandlungen 164:194–199.
- Gatesy SM. 1990. Caudefemoral musculature and the evolution of theropod locomotion.
   Paleobiology 16:170–186.
- Gatesy SM. 1991. Hind limb movements of the American alligator (*Alligator mississippiensis*) and
   postural grades. J Zool 224:577–588.
- Gatesy SM. 1994. Neuromuscular diversity in archosaur deep dorsal thigh muscles. Brain Behav Evol
   43:1–14.
- Gatesy SM. 1997. An electromyographic analysis of hindlimb function in *Alligator* during terrestrial
   locomotion. J Morphol 234:197–212.
- Gatesy SM. 1999. Guineafowl hind limb function. II: Electromyographic analysis and motor pattern
   evolution. J Morphol 240:127-142.
- Gatesy SM, Biewener AA. 1991. Bipedal locomotion: effects of speed, size and limb posture in birds
   and humans. J Zool 224:127–147.
- Gatesy SM, Dial KP. 1993. Tail muscle activity patterns in walking and flying pigeons (*Columba livia*). J
   Exp Biol 176:55-76..
- Gatesy SM, Dial KP. 1996. From frond to fan: *Archaeopteryx* and the evolution of short-tailed birds.
   Evolution 50:2037-2048.
- Gauthier J, Kluge AG, Rowe T. 1988. Amniote phylogeny and the importance of fossils. Cladistics
   4:105–209.
- Goetz JE, Derrick TR, Pedersen DR, Robinson DA, Conzemius MG, Baer TE, Brown TD. 2008. Hip joint
   contact force in the emu (*Dromaius novaehollandiae*) during normal level walking. J Biomech
   41:770–778.
- Gordon JC, Rankin JW, Daley MA. 2015. How do treadmill speed and terrain visibility influence
   neuromuscular control of guinea fowl locomotion? J Exp Biol 218:3010–3022.

- Goslow GE, Wilson D, Poore SO. 2000. Neuromuscular correlates to the evolution of flapping flight in
   birds. Brain Behav Evol 55:85–99.
- Grillner S, Wallén P. 1985. Central pattern generators for locomotion, with special reference to
   vertebrates. Annu Rev Neurosci 8:233–261.
- Hedges SB, Poling LL. 1999. A molecular phylogeny of reptiles. Science 283:998–1001.
- Herrel A, Vanhooydonck B, Porck J, Irschick DJ. 2008. Anatomical basis of differences in locomotor
  behavior in anolis lizards: A comparison between two ecomorphs. Bull Mus Comp Zool
  159:213–238.
- Higham TE, Biewener AA, Wakeling JM. 2008. Functional diversification within and between muscle
   synergists during locomotion. Biol Lett 4:41–44.
- Higham TE, Jayne BC. 2004. In vivo muscle activity in the hindlimb of the arboreal lizard, *Chamaeleo calyptratus*: general patterns and the effects of incline. J Exp Biol 207:249–261.
- Higham TE, Nelson FE. 2008. The integration of lateral gastrocnemius muscle function and
  kinematics in running turkeys. Zoology (Jena) 111:483–493.
- Hudson GE, Schreiweis DO, Wang SYC, Lancaster DA. 1972. A numerical study of the wing and leg
   muscles of tinamous Tinamidae. Northwest Science 46:207-255.
- Hutchinson J. 2001. The evolution of pelvic osteology and soft tissues on the line to extant birds
  (Neornithes). Zool J Linn Soc 131:123–168.
- Hutchinson JR. 2002. The evolution of hindlimb tendons and muscles on the line to crown-group
   birds. Comp Biochem Physiol Part A, Mol Integr Physiol 133:1051–1086.
- Hutchinson JR, Gatesy SM. 2000. Adductors, abductors, and the evolution of archosaur locomotion.
   Paleobiology 26:734–751.
- Jacobson RD, Hollyday M. 1982. A behavioral and electromyographic study of walking in the chick. J
   Neurophysiol 48:238–256.
- Jayne BC, Bennett AF, Lauder GV. 1990. Muscle recruitment during terrestrial locomotion: How
   speed and temperature affect fibre type use in a lizard. J Exp Biol.
- Jenkins FA, Goslow GE. 1983. The functional anatomy of the shoulder of the savannah monitor lizard
   (Varanus exanthematicus). J Morphol 175:195–216.
- Lamas LP, Main RP, Hutchinson JR. 2014. Ontogenetic scaling patterns and functional anatomy of the
   pelvic limb musculature in emus (*Dromaius novaehollandiae*). PeerJ 2:e716.
- Lamas LRGP. 2015. Musculoskeletal biomechanics during growth on emu (*Dromaius*; Aves): an
   integrative experimental and modelling analysis [Doctoral dissertation]. Royal Veterinary
   College, United Kingdom.
- Lyson TR, Sperling EA, Heimberg AM, Gauthier JA, King BL, Peterson KJ. 2012. MicroRNAs support a
   turtle + lizard clade. Biol Lett 8:104–107.
- Marsh RL, Ellerby DJ, Carr JA, Henry HT, Buchanan CI. 2004. Partitioning the energetics of walking
   and running: swinging the limbs is expensive. Science 303:80–83.

- Meers MB. 2003. Crocodylian forelimb musculature and its relevance to Archosauria. Anat Rec A
   Discov Mol Cell Evol Biol 274:891–916.
- Millard M, Uchida T, Seth A, Delp SL. 2013. Flexing computational muscle: modeling and simulation
   of musculotendon dynamics. J Biomech Eng 135:021005.
- Monticelli P, Ronaldson HL, Hutchinson JR, Cuff AR, d Ovidio D, Adami C. 2019. Medetomidine ketamine-sevoflurane anaesthesia in juvenile Nile crocodiles (*Crocodylus niloticus*)
   undergoing experimental surgery. Vet Anaesth Analg 46:84-89.
- Muir GD, Gosline JM, Steeves JD. 1996. Ontogeny of bipedal locomotion: walking and running in the
   chick. J Physiol (Lond) 493:589–601.
- Perreault EJ, Heckman CJ, Sandercock TG. 2003. Hill muscle model errors during movement are
   greatest within the physiologically relevant range of motor unit firing rates. J Biomech
   36:211–218.
- Prum RO, Berv JS, Dornburg A, Field DJ, Townsend JP, Lemmon EM, Lemmon AR. 2015. A
  comprehensive phylogeny of birds (Aves) using targeted next-generation DNA sequencing.
  Nature 526:569–573.
- Rankin JW, Rubenson J, Hutchinson JR. 2016. Inferring muscle functional roles of the ostrich pelvic
  limb during walking and running using computer optimization. J R Soc Interface
  13:20160035.
- Rasmussen S, Chan AK, Goslow GE. 1978. The cat step cycle: electromyographic patterns for
   hindlimb muscles during posture and unrestrained locomotion. J Morphol 155:253–269.
- Reilly SM. 1995. Quantitative electromyography and muscle function of the hind limb during
   quadrupedal running in the lizard *Sceloporus clarki*. Zoology 98:263–263.
- Reilly SM. 1998. Sprawling locomotion in the lizard *Sceloporus clarkii*: speed modulation of motor
   patterns in a walking trot. Brain Behav Evol 52:126–138.
- Reilly SM, Blob RW. 2003. Motor control of locomotor hindlimb posture in the American alligator
   (Alligator mississippiensis). J Exp Biol 206:4327–4340.
- Reilly SM, Willey JS, Biknevicius AR, Blob RW. 2005. Hindlimb function in the alligator: integrating
   movements, motor patterns, ground reaction forces and bone strain of terrestrial
   locomotion. J Exp Biol 208:993–1009.
- Rivera ARV, Blob RW. 2010. Forelimb kinematics and motor patterns of the slider turtle (*Trachemys scripta*) during swimming and walking: shared and novel strategies for meeting locomotor
   demands of water and land. J Exp Biol 213:3515–3526.
- Rivera ARV, Blob RW. 2013. Forelimb muscle function in pig-nosed turtles, *Carettochelys insculpta*:
   testing neuromotor conservation between rowing and flapping in swimming turtles. Biol Lett
   9:20130471.
- Rivera ARV, Wyneken J, Blob RW. 2011. Forelimb kinematics and motor patterns of swimming
   loggerhead sea turtles (*Caretta caretta*): are motor patterns conserved in the evolution of
   new locomotor strategies? J Exp Biol 214:3314–3323.

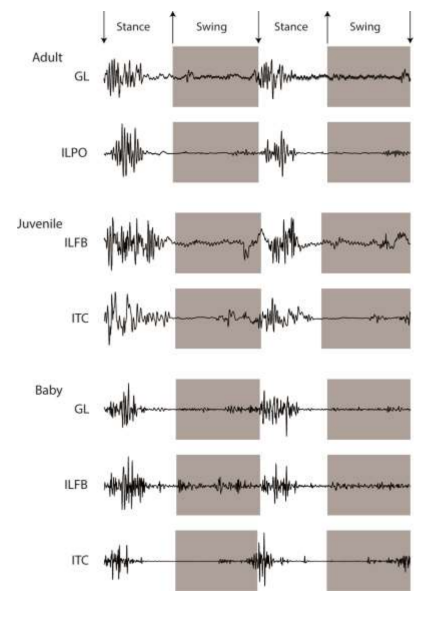
- Roberts TJ, Gabaldón AM. 2008. Interpreting muscle function from EMG: lessons learned from direct
   measurements of muscle force. Integr Comp Biol 48:312–320.
- Romer AS. 1923. Crocodilian pelvic muscles and their avian and reptilian homologues. Bulletin of the
   AMNH 48(15).
- Rowe T. 1986. Homology and evolution of the deep dorsal thigh musculature in birds and other
   reptilia. J Morphol 189:327–346.
- Schoenfuss HL, Roos JD, Rivera ARV, Blob RW. 2010. Motor patterns of distal hind limb muscles in
   walking turtles: Implications for models of limb bone loading. J Morphol 271:1527–1536.
- 812 Smith KK. 1994. Are neuromotor systems conserved in evolution? Brain Behav Evol 43:293–305.
- Tobalske BW, Jackson BE, Dial KP. 2017. Ontogeny of fight capacity and pectoralis function in a precocial ground bird (*Alectoris chukar*). Integr Comp Biol 57:217–230.
- Vanden Berge JC, Zweers GA. 1993. Myologia. In: Baumel JJ, King AS, Breazile JE, Evans HE, Vanden
  Berge JC, editors. *Handbook of Avian Anatomy: Nomina Anatomica Avium*. Cambridge,
  Massachusetts, USA: Nuttal Ornithological Club.
- Witmer LM. 1995. The extant phylogenetic bracket and the importance of reconstructing soft tissues
   in fossils. In: Thomason JJ, editor. Functional Morphology in Vertebrate Paleontology. New
   York: Cambridge University Press. p. 19–33.
- Yonezawa T, Segawa T, Mori H, Campos PF, Hongoh Y, Endo H, Akiyoshi A, Kohno N, Nishida S, Wu J,
  Jin H, Adachi J, Kishino H, Kurokawa K, Nogi Y, Tanabe H, Mukoyama H, Yoshida K,
  Rasoamiaramanana A, Yamagishi S, Hayashi Y, Yoshida A, Koike H, Akishinonomiya F,
  Willerslev E, Hasegawa M. 2017. Phylogenomics and morphology of extinct paleognaths
  reveal the origin and evolution of the ratites. Curr Biol 27:68–77.
- 826

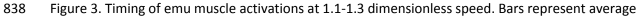
- 828 Figure 1. Avian and crocodile muscles from which EMG data were obtained . A) Avian hindlimb is
- 829 from a representative tinamou; figure modified from (Hudson et al., 1972). Muscle abbreviations
- 830 from Table 2. B) Crocodile hindlimb dorsal view, C) crocodile forelimb ventral view. Both right limbs,
- 831 modified from Allen et al., (2014). Muscle abbreviations in Table 3.



834 Figure 2. Filtered EMG signals from three emus at three ages, showing the signal variation at 1.1-1.3

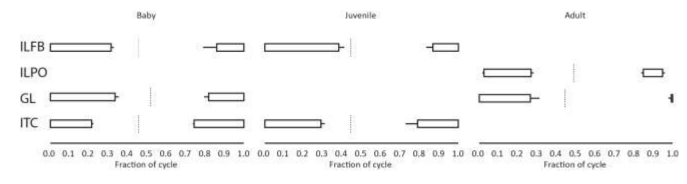
835 dimensionless speed.





839 periods of activity, with the lines representing 95% confidence intervals. Vertical dashed lines are

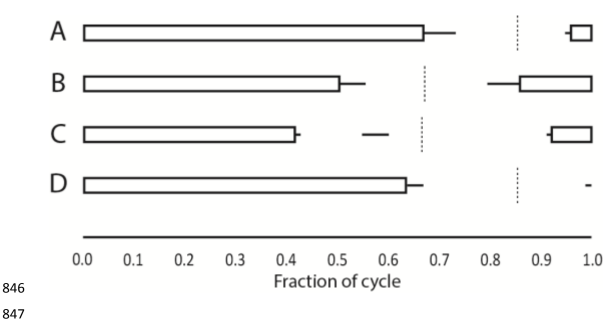
840 foot-off events.



841 842

843 Figure 4. Timing of tinamou muscle activations for A-C) GL, and D) FL muscles. A,D) *u* = 0.06, B) *u* =

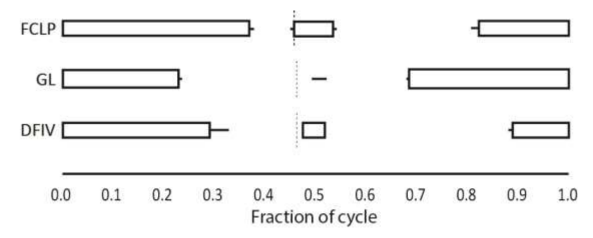
844 0.23, C) u = 0.29. Bars represent average period of activity, with the lines representing 95%



845 confidence intervals. Vertical dashed lines are foot-off events.

- 848 Figure 5. Turkey muscle activation timing at 1.2 dimensionless speed for FCLP, GL and DFIV muscles.
- Bars represent average period of activation, with the lines representing 95% confidence intervals. 849

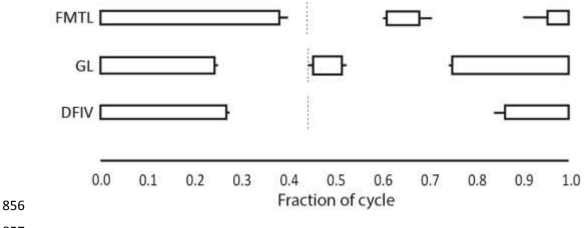
850 Vertical dashed lines are foot-off events.



851 852

853

- Figure 6. Timing of guineafowl muscle activity at 1.2 dimensionless speed for FMTL, GL and DFIV
- 854 muscles. Bars represent average activations, with the lines representing 95% confidence intervals.
- Vertical dashed lines are foot-off events. 855

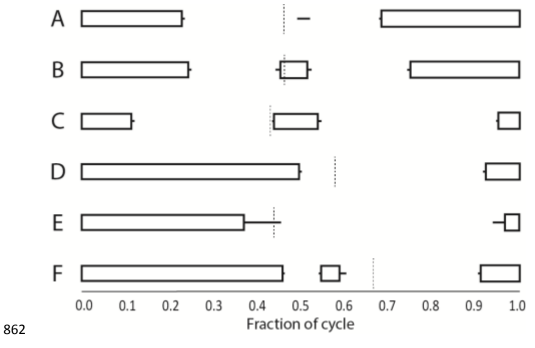


858 Figure 7. Activation timing for M. gastrocnemius pars lateralis (GL) across bird species at similar

dimensionless speed (1.1-1.3*u*). A) turkey, B) guineafowl, C) pheasant, D) quail, E) emu, F) tinamou

860 (0.29*u*). Bars represent average period of activation, with the lines representing 95% confidence

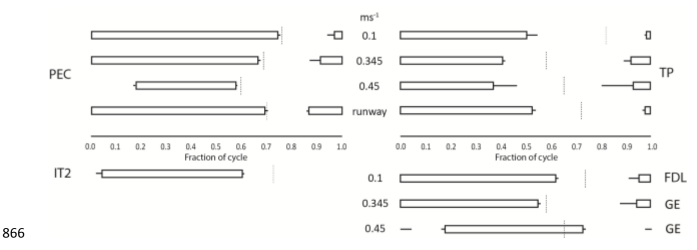
861 intervals. Vertical dashed lines are foot-off events.



863

864 Figure 8. Timing of crocodile muscle activity. Bars represent average activations, with the lines

865 representing 95% confidence intervals. Vertical dashed lines are foot-off events.



- Figure 9. Phylogeny of Sauria (based on Gauthier et al., 1988; Hedges and Poling, 1999; Field et al.,
- 2014; Prum et al., 2015), with the most parsimonious reconstruction of the evolution of EMG
- patterns mapped onto it (following Gatesy 1994, 1999). Muscle abbreviations are in Tables 2 and 3.
- 871 "?" for turtle stem reflects controversy over their precise phylogenetic position. Nodes for Sauria,
  872 Archosauria and Aves are annotated with key ancestral EMG patterns for muscles focused on in this
- study; simplified into "Stance" (circle filled on right half) for mainly stance phase activity (potentially
- with some late swing phase), "Swing" (circle filled on left half) for mainly swing phase activity, and a
- 875 "Stance" circle rotated 30 degrees anticlockwise for the more pronounced earlier swing phase
- 876 activity (and earlier stance phase end of activity) evident in the GL of Aves. Additional EMG data for
- ducks (Biewener and Corning, 2001) and pigeons (Gatesy and Dial, 1993, 1996) further bolster the
- 878 results here for Aves but for simplicity are not shown.

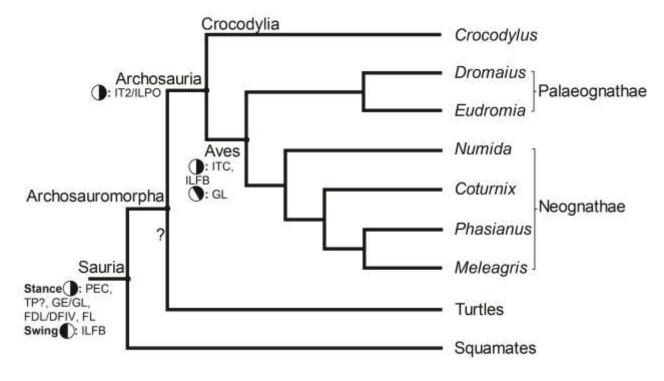


Table 1. Species used in the study

Species	Common name	Ontogenetic stage	Number of individuals	Sex	Mass (kg)
Crocodylus niloticus	Nile crocodile	Juvenile	4	F	1.59-4.63
Dromaius novaehollandiae	Emu	Baby, Juvenile, Adult	2 2 2	Unknown	3.95-4.1, 17-18, 36-37
Eudromia elegans	Elegant- crested tinamou	Adult	2	F,M	0.555-0.616
Numida meleagris	Guineafowl	Adult	2	F,M	1.5, 1.7
Phasianus colchicus	Common pheasant	Adult	2	М	1.0, 1.1
Coturnix virginianus	Bobwhite quail	Adult	2	F	0.15, 0.17
Meleagris gallopavo	Wild turkey	Adult	2	F	5.2, 6.0

Table 2. Bird muscles that EMG data were obtained from, with previously reported anatomy and function. Nomenclature follows Vanden Berge and Zweers (1993) for Aves. "Actions" (presumed potential functions around joints crossed) are inferred from anatomy following Vanden Berge and Zweers (1993) and Lamas et al. (2014).

Muscle	Origin	Insertion	Action	
lliotrochantericus caudalis (ITC)	Preacetabular ilium	Trochanteric crest of proximal femur	Hip flexor, abductor and internal rotator	
Iliotibialis lateralis pars postacetabularis (ILPO)	Dorsal postacetabular ilium	Patella and cranial tibial crest of tibiotarsus	Hip extensor, abductor and external rotator; knee extensor	
Iliofibularis (ILFB)	Postacetabular ilium	Fibula (M. iliofibularis tubercle)	Hip extensor, abductor and external rotator; knee flexor	
Flexor cruris lateralis pars pelvica (FCLP)	Posterior rim of terminal iliac process	GE/GL and flexor cruris medialis tendon	Hip extensor, abductor and external rotator; knee flexor	
Femorotibialis lateralis (FMTL)	Lateral surface of femur	Craniolateral surface of proximal tibiotarsus	Knee extensor	
Gastrocnemius pars lateralis (GL)	Caudal side of lateral condyle of femur	Caudal surface of proximal (tarso)metatarsus (i.e. hypotarsus of birds)	Knee flexor; ankle extensor	
Fibularis longus (FL)	-		Ankle extensor (and potentially digital flexor)	
Flexor perforatus digiti IV (DFIV)			Ankle extensor; digit IV flexor	

Table 3. Crocodile muscles that EMG data were obtained from, with previously reported anatomy and function. Nomenclature follows Meers (2003) and Allen et al. (2014) and references therein for Crocodylia. "Actions" (presumed potential functions around joints crossed) are inferred from anatomy.

Muscle	Origin	Insertion	Action		
Pectoralis (PEC)	Ventral surface of sternum, ribcage, surrounding area	Deltopectoral crest of humerus	Glenohumeral extensor, abductor and supinator		
Transversus perinei (TP)	Ischium	Centra of caudal vertebrae 1+2	Unknown: possibly shaping tail base		
lliotibialis 2 (IT2)	Dorsal ilium	Cranial side of proximal tibia	Hip abductor (also long-axis rotator and flexor/extensor?); knee extensor		
Gastrocnemius externus (GE)	Caudal side of lateral condyle of femur	Caudal surface of proximal (tarso)metatarsus	Knee flexor; ankle extensor		
Flexor digitorum longus – hindlimb (FDL)	Disto-lateral femoral condyle	Distal pes	Digital flexor; ankle extensor		

Species		Tina	mou			Turkey		Pheasant	Quail		Guineafow	-
Muscle		GL		FL	GL	FCLP	DFIV	GL	GL	GL	DFIV	FMTL
N individuals	1	1	1	1	2	2	2	1	2	2	2	2
Surface	Treadmill											
Speed (ms-1)	0.10	0.35	0.45	0.10	2.15	2.15	2.15	1.7	1.12	1.7	1.7	1.7
u	0.06	0.23	0.29	0.06	1.27	1.27	1.27	1.25	1.23	1.28	1.28	1.28
# trials	2	2	2	2	5	5	5	1	6	6	6	6
# steps	7	7	17	7	46	46	46	14	45	54	54	54

# Table 4. EMG data summary for bird subjects.

Table 5. EMG trial and data summary for Nile crocodile subjects.

Species		Nile crocodile													
Muscle		Т	P		IT2	FDL	PEC GL					iL			
N individuals	1 2 1 1				1	1	1 1 1 1			1	1				
Surface	Runway	Treadmill	Treadmill	Treadmill	Treadmill	Treadmill	Runway	Treadmill	Treadmill	Treadmill	Treadmill	Treadmill			
Speed (ms <sup>-1</sup> )	0.10	0.10	0.35	0.45	0.10	0.10	0.10	0.10	0.35	0.45	0.35	0.45			
u	0.07	0.07	0.26	0.33	0.07	0.07	0.07	0.07	0.26	0.33	0.26	0.33			
# trials	5	9	2	2	2	5	5	4	2	2	2	2			
# steps	14	25	8	7	6	16	14	19	8	9	8	7			

	Emu - adult											
Muscle			GL			ILPO						
N individuals	1	1	2	2	2	1	1	1	1	1		
Surface	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway		
Speed (ms <sup>-1</sup> )	0.98-1.4	1.4-2.02	2.02-2.65	2.65-3.23	3.23-3.7	0.98-1.4	1.4-2.02	2.02-2.65	2.65-3.23	3.23-3.7		
u	0.3-0.5	0.5-0.7	0.7-0.9	0.9-1.1	1.1-1.3	0.3-0.5	0.5-0.7	0.7-0.9	0.9-1.1	1.1-1.3		
# trials	2	4	12	21	7	2	4	6	8	3		
# steps	4	8	24	41	13	4	8	12	16	6		

	Emu - juvenile												
Muscle	IL	FB		ITC									
N individuals	1	1	1	1	1	1	1	1					
Surface	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway					
Speed (ms <sup>-1</sup> )	2.9-3.3	3.3-4.0	1.19-1.3	1.4-1.9	1.9-2.4	2.4-2.9	2.9-3.3	3.3-4.0					
u	1.1-1.3	1.3-1.5	0.3-0.5	0.5-0.7	0.7-0.9	0.9-1.1	1.1-1.3	1.3-1.5					
# trials	10	9	2	1	5	7	10	9					
# steps	18	16	4	2	11	15	18	16					

	Emu - baby															
Muscle			G	iL			ILFB				ITC					
N individuals	1 1 2 1 2 2					2	1	1	1	1	1	1	2	1	1	1
Surface	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway	Runway
Speed (ms <sup>-1</sup> )	0.6-0.92	0.92-1.3	1.3-1.7	1.7-2.0	2.0-2.4	2.4-2.7	1.3-1.7	1.7-2.0	2.0-2.4	2.4-2.7	0.6-0.92	0.92-1.3	1.3-1.7	1.7-2.0	2.0-2.4	2.4-2.7
u	0.3-0.5	0.5-0.7	0.7-0.9	0.9-1.1	1.1-1.3	1.3-1.5	0.7-0.9	0.9-1.1	1.1-1.3	1.3-1.5	0.3-0.5	0.5-0.7	0.7-0.9	0.9-1.1	1.1-1.3	1.3-1.5
# trials	2	3	11	8	19	10	7	8	9	1	2	1	9	2	10	9
# steps	6	10	43	34	81	43	26	34	40	4	6	3	31	9	41	39

Table 6. EMG trial and data summary for emu subjects across ontogeny.

Table 7. Cross-correlation analysis results for lateral gastrocnemius (GL/GE) EMG timings for all species at 1.1-1.3*u* or maximal speed obtained. "Max Cor" is the maximum correlation between the average signals; "offset" is the offset of the maximum correlations between the species (as fraction of a stride). "Croc" is Nile crocodile; "Guinea" is guineafowl.

	Croc	Emu	Quail	Tinamou	Turkey	Guinea	Pheasant	
Croc	1.000	0.855	0.802	0.817	0.487	0.478	0.746	Max Cor
CIUC	0.000	0.178	0.230	0.228	0.266	0.230	0.183	Offset
Emu		1.000	0.940	0.941	0.728	0.699	0.834	Max Cor
LIIIU		0.000	0.000	0.000	0.014	0.020	0.000	Offset
Quail			1.000	0.953	0.814	0.789	0.900	Max Cor
Quali			0.000	0.000	0.004	0.012	0.000	Offset
Tinamou				1.000	0.759	0.753	0.869	Max Cor
Tillalliou				0.000	0.000	0.000	0.000	Offset
Turkey					1.000	0.970	0.788	Max Cor
Turkey					0.000	0.000	0.000	Offset
Guinea						1.000	0.782	Max Cor
Guillea						0.000	0.000	Offset