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8 1 Scaling of primate forearm muscle architecture as it relates to locomotion and posture
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2
3 70 ABSTRACT
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6 71 It has been previously proposed that distal humerus morphology may reflect the
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8 72 locomotor pattern and substrate preferred by different primates. However,
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10 73 relationships between these behaviors and the morphological capabilities of muscles
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12 74 originating on these osteological structures have not been fully explored. Here, we
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14 75 present data about forearm muscle architecture in a sample of 44 primate species (n =
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16 76 55 specimens): 9 strepsirrhines, 15 platyrrhines, and 20 catarrhines. The sample
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18 77 includes all major locomotor and substrate use groups. We isolated each antebrachial
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20 78 muscle and categorized them into functional groups: wrist and digital extensors and
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22 79 flexors, antebrachial mm. that do not cross the wrist, and functional combinations
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24 80 thereof. Muscle mass, physiological cross-sectional area (PCSA), reduced PCSA (RPCSA),
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26 81 and fiber length (FL) are examined in the context of higher taxonomic group, as well as
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28 82 locomotor/postural and substrate preferences. Results show that muscle masses, PCSA,
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30 83 and RPCSA scale with positive allometry while FL scales with isometry indicating that
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32 84 larger primates have relatively stronger, but neither faster nor more flexible, forearms
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34 85 across the sample. When accounting for variation in body size, we found no statistically
35
36 86 significant difference in architecture among higher taxonomic groups or
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38 87 locomotor/postural groups. However, we found that arboreal primates have
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40 88 significantly greater FL than terrestrial ones, suggesting that these species are adapted
41
42 89 for greater speed and/or flexibility in the trees. These data may affect our
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44 90 interpretation of the mechanisms for variation in humeral morphology and provide
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46 91 information for refining biomechanical models of joint stress and movement in extant
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92 and fossil primates.

93 Key words: primate, forearm, locomotion, arboreal, terrestrial, muscle architecture

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10 Multiple studies have suggested that osteological differences in the distal
11 humerus among primates may reflect variation in the force production capabilities of
12 the forearm musculature, likely necessitated by differences in substrate interaction
13 between primates practicing different locomotor patterns (Jenkins, 1973; Ziemer, 1977;
14 Szalay and Dagosto, 1980; Feldesman, 1982; Fleagle and Simons, 1982; Fleagle and
15 Meldrum, 1988; Harrison, 1989; Frost and Delson, 2002). However, in addition to
16 osteological features associated with mechanical advantage, muscle strength may be
17 affected by a number of variables, including muscle mass and muscle fiber architecture.
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28 Jenkins (1973) describes a condition by which torsional stresses at the elbow
29 joint are produced by the force of the extrinsic digital and wrist flexors during pronated
30 hand postures. According to his model (Jenkins, 1973), this force can be
31 counterbalanced in three ways: 1) reducing the flexor muscle mass, 2) increasing the
32 extensor muscle mass, or 3) shifting the flexor muscles posteriorly to run behind the
33 elbow joint. This elegant explanation has been highly cited in the literature and appears
34 to correspond well with the observed skeletal morphology of cercopithecoid primates
35 (Napier and Davis, 1959; Birchette, 1982; Fleagle and Simons, 1982; Harrison, 1989;
36 McCrossin et al., 1998; Frost and Delson, 2002). Although analyses of extrinsic forearm
37 muscle weights have been performed for hominoids and some catarrhine monkeys
38 (Tuttle, 1969; Tuttle, 1972), there is little information available for platyrrhine and
39 strepsirrhine primates.
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4 136 While the argument that large medial epicondyles support large flexor muscles is
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7 137 clearly logical, muscle force is not reliably predicted by muscle mass or volume (Gans
8
9 138 and Bock, 1965). Rather, muscle action is affected by muscle architecture, including
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12 139 fiber length and degree of pennation. The use of physiological cross-sectional area
13
14 140 (PCSA) and reduced physiological cross-sectional area (RPCSA) takes these variables into
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17 141 account and provides a better estimate of maximum muscle force than weight
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19 142 measurements alone (Payne et al., 2006; Taylor and Vinyard, 2008). Theoretically, a
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22 143 given muscle compartment of an animal could produce more force with less total
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24 144 volume by altering muscle architecture. Subtle differences in muscle architecture
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27 145 introduce the possibility that the bony area of muscular insertion does not reflect the
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29 146 muscle mass, but instead reflects the force produced by that muscle. Tuttle (1972) was
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32 147 aware of this fact and avoided the use of absolute muscle mass and/or volume in his
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34 148 study of anthropoid hands and feet.

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37 149 The model proposed by Jenkins (1973) implies that the frequency of use of a
38
39 150 particular muscle group in locomotion is reflected in variable volume of the musculature
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42 151 and its osteological origin (i.e., greater employment in locomotion = bigger muscle
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44 152 volume = larger bony insertion). Following this model, we would expect animals
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47 153 involved in locomotor categories employing frequent gripping (i.e., arboreal
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49 154 quadrupedalism, climbing, and suspension) to require greater force of the digital flexors
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52 155 compared to animals that are predominantly terrestrial. Terrestrial animals would be
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54 156 expected to require generation of large forces by their deep digital flexors (in toe-off)
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57 157 and their pronators (in quadrupedalism), but less force in their wrist and superficial
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3 158 digital flexors. The model predicts that greater force will be generated by either
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6 159 increased muscle volume of the appropriate groups, or an appropriate change in
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9 160 muscular architecture. Further, any increase in muscle volume is expected to generate
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11 161 a larger bony origin site (greater epicondylar projection).
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16 163 Muscle fiber architecture and its relationship to posture, substrate use and locomotor
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18 164 patterns
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23 166 In our previous studies (Perry and Hartstone-Rose, 2010; Hartstone-Rose and
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26 167 Perry, 2011; Hartstone-Rose et al., 2012b; Hartstone-Rose et al., 2015) the muscle
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28 168 architecture of the masticatory system in felids and lemurs co-vary with the mechanical
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31 169 requirements of differing diets—specifically with gape and bite force. In both clades,
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34 170 the length of the masticatory muscle fibers seems to be adapted for the size of food
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36 171 items, while force variables scale isometrically with body size. In other words, the mass
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38 172 and physiological cross-sectional area of the masticatory muscles scale tightly with body
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41 173 mass, but provide no real dietary behavioral signals. Muscle fiber length, on the other
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43 174 hand, does reveal information about dietary behavior. Because of this common pattern
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46 175 in the masticatory muscles, we suspect that muscle fiber architecture plays an
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48 176 important role in other anatomical regions, for instance as an indicator of adaptation in
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51 177 the muscles of locomotion. An examination of muscle fiber architecture in the limb
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53 178 muscles may indicate whether limb muscles are adapted for strength or speed—the
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56 179 latter, for instance, may be of optimum advantage in aid of vertical leaping. We might
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4 180 also find correlations between muscle fiber structure and substrate use (e.g., terrestrial
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7 181 vs. arboreal) as arboreal primates rely more heavily on quick movement through the
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9 182 trees than terrestrial primates do on land, or higher taxonomic group (between
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11 183 catarrhines, platyrrhines, and strepsirrhines) since selection or drift could drive
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13 184 morphological changes in muscle fiber architecture as well. Fiber type analysis would
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15 185 add even greater depth to this type of analysis, but is beyond the scope of the current
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17 186 study.
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24 188 Muscle fiber architecture
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29 190 Skeletal muscle fiber structure should vary across skeletal regions and across
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32 191 taxa in patterns that are related to the selective pressures of that anatomical region –
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34 192 for instance those imposed on the muscles of mastication by specific diets and those
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36 193 imposed on the forelimb muscles by specific locomotor behaviors. Skeletal muscles can
37
38 194 be seen as a collection of fibers arranged in parallel. While these fibers have a relatively
39
40 195 consistent diameter in muscles of varying sizes, the spatial arrangement of fibers can
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42 196 vary significantly. It is the arrangement of these fibers that is responsible for muscle
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44 197 contraction and force production (Brinckmann, 2002). Muscle fibers may be oriented
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46 198 longitudinally to the force generating axis of the muscle, or they may be pennate
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48 199 (oriented at an angle). In muscles of equivalent gross dimensions, muscles with
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50 200 longitudinally-oriented fibers are longer, and thus the muscle has greater potential to
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52 201 stretch (since stretch is proportional to the resting length of the muscle) and also to
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3 202 move more quickly (since each muscle fiber theoretically contracts at the same rate and
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6 203 longer fibers cover more distance in that same amount of time). Pennation serves to
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9 204 increase the number of muscle fibers with in a given muscle volume. Fibers have a
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11 205 relatively consistent cross section, and cross-sectional area is directly proportional to
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13 206 force production. Pennation increases muscle strength as a consequence of this
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16 207 increase in fiber number. However, pennate muscles have shorter fiber lengths for the
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19 208 same muscle volume. Thus, a fundamental tradeoff exists in muscle fiber architecture:
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21 209 for a given muscle volume, muscles can be adapted for stretch and speed (fewer longer
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23 210 fibers), or adapted for strength (greater number of shorter fibers). If a muscle needs to
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26 211 be both strong and fast and/or flexible, then the muscle must be of greater volume than
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29 212 one that is adapted for only one of these states, or for a compromise thereof (Anapol
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31 213 and Barry, 1996).

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33 214 As a consequence of differences in muscle fiber orientation, the *anatomical* cross-
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36 215 sectional area of a muscle is not always proportional to its force generating capacity.
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39 216 The *physiological* cross-sectional area (PCSA) of a muscle – the cross-sectional area
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41 217 measured perpendicular to the longitudinal axis of fiber orientation – is a more accurate
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43 218 measure of muscle force production because it represents the sum of the individual
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46 219 muscle fiber cross-sectional areas (Maughan et al., 1983; Brinckmann, 2002). *Reduced*
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49 220 physiological cross-sectional area (RPCSA) – another measure of muscle force – takes
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51 221 into account the amount of force lost due to the angle of muscle fibers in a pennate
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54 222 muscle that runs orthogonal to the muscle's line of action – essentially by removing the
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56 223 vector of force imparted by pennation that is perpendicular to the total muscle's line of
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224 action (Anapol and Barry, 1996; Perry et al., 2011).

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226 Locomotor Pattern

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228 The primate order is characterized by a great diversity of locomotor behaviors
229 (Hunt et al., 1996; Fleagle, 1999). Field research has established that greater variation in
230 locomotion exists among primates than was originally thought, both between and
231 within species. Since the early attempts to classify primate locomotion into categories
232 (Prost, 1965a, 1965b) it has been clear that such a task is quite difficult, though
233 anatomists and especially students who use biomechanical approaches have long
234 recognized that a complete understanding of primate musculoskeletal system must rely
235 especially on quantitative primate positional behavior data (Hunt et al., 1996). The aim
236 here is to provide only broad locomotor distinctions, recognizing that some variability in
237 locomotor/postural behavior exists within the taxonomic groups shown in Table 1. As
238 described in the methods section below, we have assigned each species represented in
239 our sample to locomotion/posture and substrate use categories based on the available
240 behavioral literature.

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242 HYPOTHESES AND PREDICTIONS

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244 The primary goal of this study is to evaluate the potential correlation between
245 extrinsic hand/wrist muscle architecture and locomotor patterns/posture, substrate

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3 246 use, and higher taxonomic group across the Order Primates. We explore the scaling
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6 247 patterns of individual architectural variables with body size, in an attempt to elucidate
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9 248 whether there are differences in allometric scaling patterns associated with contrasts in
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11 249 higher taxonomic group, locomotion/posture, and substrate categories. We also test
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13 250 predictions related to the distribution of relative muscle volume and muscle among
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16 251 these groups—i.e. do statistically significant differences occur among primates of
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18 252 differing higher taxonomic group, locomotion/posture, and substrate preferences, when
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21 253 variation related to body size is taken into account?
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26 255 Our hypotheses regarding forearm muscle architecture and the behaviors in question
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28 256 are as follows:
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33 258 1) Because of their reliance on speed to move through their habitat, arboreal primates
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36 259 will have relatively greater forearm flexor fiber length (FL) compared to terrestrial
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38 260 primates. Alternatively, because of their reliance on grip strength to hang onto trees,
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41 261 arboreal primates will have relatively high PCSA and RPCSA compared to terrestrial
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43 262 primates.
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48 264 2) Vertical clingers and leapers will, because of their reliance on quick movement
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51 265 through the trees, have relatively greater forearm FL compared to quadrupedal and
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53 266 suspensory primates. Alternatively, because of their more obtuse wrist position, vertical
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56 267 clinging and leaping primates will have relatively shorter forearm FL compared to
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268 quadrupedal and suspensory primates.

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270 How these variables interact with higher taxonomic group (to examine the influence of
271 founder/drift effects) and across body size (to examine allometric effects) will also be
272 assessed.

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274 MATERIALS AND METHODS

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276 The specimens in this study include ten species of strepsirrhines, fifteen species
277 of platyrrhines, and twenty species of catarrhines (Table 1). More than one individual of
278 eleven species was dissected (*Eulemur fulvus*, *Aotus azarae*, *Callithrix jacchus*, *Callithrix*
279 *geoffroyi*, *Cebuella pygmaea*, *Saimiri sciureus*, *Sapajus apella*, *Chlorocebus aethiops*,
280 *Miopithecus talapoin*, *Cercocebus atys*, *Gorilla gorilla gorilla*); in these cases the masses
281 of the individuals dissected were averaged, as well as all functional values. If the
282 individual's weight was unavailable, the average for its sex was taken from Fleagle
283 (1999)—if the sex was unknown, a species average was taken from the same reference.
284 All were adult animals and (except the aye-aye, *Daubentonia madagascariensis*) were
285 from captive facilities in the United States and Spain, with the majority of specimens
286 being from Spanish zoos and dissected at the Universidad de Valladolid. The aye-aye
287 was a wild born specimen that died at Parc Tsimbazaza, Antananarivo Madagascar. The
288 same specimen was used for a previous study on the masticatory muscle structure

289 (Perry et al., 2013). All specimens were fresh frozen with minimal degradation of the
 290 muscle tissue.

291 All antebrachial muscles were dissected from one forearm of each specimen.

292 Each muscle was isolated, removed from its insertion points on the bone, and measured
 293 for volume and muscle architecture variables. Data for each muscle was considered
 294 both individually and as a part of a broader functional group—i.e. categorized into the
 295 broad group of “flexors” (flexor carpi radialis, palmaris longus, flexor carpi ulnaris, flexor
 296 digitorum superficialis, flexor digitorum profundus, flexor pollicis longus), “extensors”
 297 (extensor carpi radialis longus, extensor carpi radialis brevis, extensor digitorum,
 298 extensor carpi ulnaris, abductor pollicis longus, extensor digiti minimi, extensor indicis,
 299 extensor pollicis longus, extensor pollicis brevis) and “others” – muscles that do not flex
 300 or extend the wrist (brachioradialis, pronator teres, pronator quadratus, supinator).

301 Subgroupings of these (e.g., wrist and digital flexors and extensors) were also analyzed.

302 Some of these specific muscles, particularly those involving fine motor control, were not
 303 distinctly identifiable in some specimens. In contrast, some species had notable
 304 elaborations and subdivisions of these muscles; for instance, the aye-aye (*Daubentonia*)
 305 had a distinct muscle that extended its third digit. It is not surprising that this unique
 306 “tap forager” has a unique muscle configuration. Since we analyzed muscles in broader
 307 groups (e.g., all digital extensors combined) and not in terms of individual identifiable
 308 muscles, these variations did not preclude analyses across the whole sample; all species-
 309 specific muscle variants were easily combined into their functional groupings.

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4 310 For those muscles crossing the radiocarpal joint, the tendons were uniformly cut
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7 311 at the level of the flexor and extensor retinacula, rather than excised from their
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9 312 insertion point within the hand. The external tendon was cut from the muscle at the
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11 313 level in which no muscle fibers continued to inset upon it (Stern, 1971). The muscle
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13 314 belly length, width, and thickness were then recorded with respect to their anatomical
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15 315 placement within the forearm, each measured to the nearest 0.01mm using digital
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17 316 calipers. The muscle bellies were then weighed to 0.0001g using a digital scale.
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22 317 Muscle fiber length was measured using a protocol modified from Rayne and
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24 318 Crawford (1972). Each muscle was immersed in 10% sulfuric acid (for all but the aye-
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26 319 aye) and cooked in an oven at 60°C. Cooking time varied between 45 minutes and 6
27
28 320 hours depending on muscle size and the amount of connective tissue present. The aye-
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30 321 aye (*Daubentonia*) was opportunistically dissected on site in Madagascar, where sulfuric
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32 322 acid was not readily available; instead, its muscles were cooked over longer periods
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34 323 (~1.5 times longer) in white vinegar (~acetic acid). This substitution was validated prior
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36 324 to using it on this specimen, and found to yield equivalent results. The acid cooking
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38 325 process dissolves the collagenous connective tissue and allows individual fascicles to be
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40 326 carefully manipulated with forceps without breakage. The fascicle lengths are then
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42 327 measured using digital calipers. An average of 40 representative fascicle bundles were
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44 328 measured per muscle, ensuring that all regions of the muscle were represented. The
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46 329 mean of these measurements was taken as the fascicle/fiber length (FL) for that muscle.
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53 330 Muscle mass (MM) and FL are used to calculate PCSA for each muscle from the
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55 331 following equation using a formula modified from Schumacher (1961):
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$$q = m/lp$$

Here, q is PCSA, m is muscle mass, l is mean fiber length, and p is a constant: the specific density of muscle. The p value used was 1.0564 g/cm^3 (Murphy and Beardsley, 1974). PCSA is in cm^2 , muscle mass is in grams (g), and fiber length is in centimeters (cm). Calculations were made after converting FL from millimeters to centimeters.

Muscle thickness, measured perpendicular to the direction of pull of the muscle (bony origin to bony insertion) in the coronal plane, is used for calculating the pennation angle (θ) to then calculate RPCSA. The following formula is used for determining θ (Anapol and Barry, 1996), in which a is muscle thickness and l is mean fiber length:

$$\sin\theta = a/l$$

Reduced physiological cross-sectional area (RPCSA) provides information about the portion of muscle cross section that applies force parallel to the muscle's overall line of action. The equation for calculating RPCSA includes pennation angle (θ) and, as its name implies, RPCSA is always smaller than PCSA unless all of the muscle fibers run parallel to the long axis of the muscle – that is, the muscle has no pennation at all. We again follow the method of Anapol and Berry (1996) in using the following equation:

$$q_r = m(\cos\theta)/lp$$

Muscle data were separated into functional groups: all flexors, all extensors, wrist and digital flexors, wrist and digital extensors, others (pronators, supinator, and brachioradialis), and all muscles combined. MM, PCSA, and RPCSA were calculated as the sum of those values for all muscles in a particular group for an individual specimen. A weighted average FL for each muscle group was calculated by adding the products of

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354 each muscle's FL by its MM and then dividing that by the total MM for the group. Prior
355 to regression of logged variables, the cube root of the cubic variables (masses) and the
356 square root of squared variables (PCSA and RPCSA) was taken so that all predicted
357 regression slopes for logged data would be equal to 1.

358 Predominant locomotion, posture and substrate use were derived from the
359 literature (Table 1). We grouped the species included in this study into three broad
360 locomotor categories: "quadrupedal," which includes primates (both arboreal and
361 terrestrial) that move mainly using a quadrupedal gait; "vertical clinging/leaping," which
362 includes primates that exhibit leaping behavior as a substantial portion of their
363 locomotion; and "suspensory," which includes primates that rely mainly on their
364 forelimbs for locomotion in the trees. We grouped both strepsirrhines (like *Hapalemur*
365 *griseus* and *Galago senegalensis*) and leaping anthropoids (callitrichines) in the vertical
366 clinging and leaping (VCL) category following Kinzey et al. (1975) and Garber (1992).

367 Many primates, though primarily arboreal, spend some time on the ground
368 foraging and moving from one foraging site to the next, while others are mainly
369 terrestrial and climb up trees only to escape predators or to sleep (Fleagle, 1999).
370 However, the generally adopted subdivision between arboreal and terrestrial primates
371 does not fully take into account the complexity of substrate use patterns. The degree of
372 terrestriality of different primates can change considerably, even among species that
373 are closely related (Gebo, 1987; McGraw, 1998, 2000). We recognize that many species
374 of guenons, for example, spend different amounts of time on the ground (Gebo and
375 Sargis, 1994); however, for the purpose of this study we classified each species as either

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3 376 arboreal or terrestrial, with terrestrial species spending the majority of their feeding
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6 377 time on the ground (Marchi et al., this issue). In the future, the inclusion of more species
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9 378 in studies of this nature could allow for more specific categorizations concerning
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11 379 substrate use.

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13 380 In order to test the prediction that there are differences in forearm muscle
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16 381 architecture across primates based on higher taxonomic group, locomotor/posture
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18 382 pattern, and substrate use, scaling patterns were evaluated using Reduced Major Axis
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21 383 (RMA) regressions of each functional variable (MM, FL, PCSA, RPCSA, organized into
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23 384 their functional groups) on the independent variable of body mass. RMA regression was
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26 385 applied across the entire primate sample, and within-group scaling patterns were
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28 386 assessed via RMA of subsets of data parsed by taxonomic group, locomotor category,
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31 387 and substrate use.

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33 388 Residuals were calculated as the perpendicular distance from the RMA
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36 389 regression line in log space. Multiple pairwise comparisons were applied to evaluate
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38 390 statistical differences in these “size-adjusted” muscle masses among taxonomic,
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41 391 locomotor/postural, and substrate-use groups. All analyses were done in JMP13 (SAS),
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43 392 using a significance criterion of $\alpha < 0.05$.

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47 48 394 RESULTS

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53 396 Allometry across the whole sample

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56 397 All muscle masses correlate strongly with body mass ($r^2 \geq 0.96$). Across the
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entire sample, muscle mass scales with body mass via significant, albeit often weak, positive allometry (confidence intervals for slope > 1.0) for the following muscle groups: total flexors + extensors, flexors, total forearm muscles, other muscles, wrist flexors (Table 2). Based on the allometry of the slopes, we can infer that larger primates have relatively larger muscle mass overall than smaller primates. Interestingly, the extensor muscle groups by themselves scale with body mass at a slope statistically indistinguishable from isometry (95% CI overlaps 1.0).

PCSA scales with positive allometry (slopes range from 1.13 to 1.47, see Table 2) with body mass for all muscle groups, with the exception of wrist extensors (WE), which trend weakly towards positive allometry (slope = 1.13, 95%CI = 0.98 to 1.29). Judging from this, it is clear that larger primates have relatively stronger forearm muscles overall, and in particular, relatively stronger forearm flexors than smaller primates. A similar scaling relationship occurs between RPCSA and body mass; however, in this case WE scales with weak positive allometry (slope = 1.13, 95% CI = 1.00 to 1.27) on body mass. Thus, even when “correcting” for pennation, larger primates’ forearms are still relatively stronger than those of smaller primates. FL scales isometrically with body mass across every muscle group, indicating that larger primates would not be expected to have relatively greater fiber lengths in their forearm muscles than smaller primates. Although there is a scaling relationship with the strength variables (larger animals are relatively stronger) there is not one with the speed/flexibility variable (FL).

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The same trends hold true when looking at the scaling of specific divisions of the

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3 420 muscles across the primate sample. That is, PCSA and RPCSA scale with positive
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5 421 allometry (slopes range from 1.13 to 1.31), while FL scales with isometry tending toward
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7 422 weak, but insignificant (95% CI for slope encompasses 1.00) negative allometry for the
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9 423 flexors and extensors when analyzed both as groups, and when separated into digital
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11 424 and wrist subdivisions. While MM for the combined flexors and extensors was
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13 425 significantly positively allometric, isometric scaling cannot be statistically excluded for
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15 426 the extensor group or the digital or wrist extensor subgroup muscle masses. As with the
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17 427 combined flexor and extensor statistics, larger primates have relatively larger PCSA and
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19 428 RPCSA (signifying greater strength) across all separate forearm functional groups.
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21 429 Likewise, they also have isometric FL trending toward negative allometry for each
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23 430 functional group. However, while the combined flexor and extensor muscles tend to be
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25 431 relatively larger (greater MM) in larger primates, this is driven only by relatively larger
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27 432 flexor MM – the MM of the extensors, although trending toward positive allometry, are
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29 433 statistically isometric.
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33 435 Analyses of the residuals of architectural variables by higher taxonomic and functional
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438 No significant differences were found among higher taxonomic groups
439 (strepsirrhines, platyrrhines and catarrhines) in size-adjusted variables (residuals from
440 RMA regression on log body mass). Thus, in our sample, none of these taxonomic
441 groups is any more or less strong, fast, or flexible in any of their forearm muscle groups.

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442 Contrary to our hypotheses, there were also no statistical differences in the
443 forearm architectural variables between the locomotion/posture groups. That is, there
444 are no statistical differences in the strength, flexibility or speed abilities of quadrupedal
445 vs. suspensory vs. vertical clinging and leaping (VCL) taxa or the quadrupeds vs. the
446 combined group of suspensory and VCL species. Also, surprisingly, the aye-aye fell
447 within the range of similarly sized quadrupedal primates across all of the architectural
448 variables despite its highly specialized manual abilities.

449 Unlike the postural/locomotor comparisons, analyses show that arboreal and
450 terrestrial species are significantly different in PCSA and RPCSA (Table 3): Although there
451 are not significant differences in any of the MM variables between arboreal and
452 terrestrial taxa (i.e., both have equally massive forearm muscles), there is some
453 evidence that terrestrial species have relatively higher PCSA and RPCSA. Although only a
454 few of these strength variables are significant or approach significance, it does appear
455 that terrestrial primates are, in some ways, stronger than arboreal species. The more
456 compelling trend is clearly in the FL data (Fig. 1): arboreal primates have relatively
457 longer forearm muscle fibers than terrestrial primates in their combined flexors and
458 extensors, and also in the sum of all forearm muscles. Most of the other subgroups of
459 muscles either reach or approach significance as well, but the unambiguously significant
460 difference in the most combined categories of muscles suggests that there really is an
461 overarching difference in the FL between terrestrial and arboreal primates – the latter
462 clearly have either more flexible or faster forearm abilities.

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3 464 DISCUSSION
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6 465 This study sought to determine whether there are significant relationships
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8 466 between locomotor/postural or substrate groups and forearm muscle architectural
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10 467 variables in primates. We examined a large number of variables grouped by wrist and
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12 468 digital flexors and extensors, “other” muscles (a grouping of the antebrachial muscles
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14 469 that do not affect the wrist or digits) and various combinations thereof. We also sought
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16 470 to use these architectural variables to identify whether specific lineages or functional
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18 471 groups of primates appear to be adapted for strength or speed/flexibility. As we have
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20 472 found in our previous studies of the masticatory apparatus (Hartstone-Rose and Perry
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22 473 2011; Hartstone-Rose et al. 2012; Perry and Hartstone-Rose 2010) and our similar
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24 474 analysis of leg muscles (Marchi et al., this issue), muscle fiber architecture clearly
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26 475 demonstrates the functional trade-off between strength and stretch with notable
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28 476 behavioral correlates.
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36 477 As we expected based on previous studies (and basic principles of allometry), our
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38 478 regressions indicate that larger primates have relatively larger and stronger forearm
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40 479 muscles than do smaller primates; when the forearm muscles were analyzed in their
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42 480 functional compartments, PCSA and RPCSA scaled with positive allometry (or with
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44 481 isometry trending toward positive allometry) with body mass in nearly every muscle
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46 482 category. Fiber length, however, scales with isometry with body mass (though this is
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48 483 not always the case with other animals; e.g. Herrel et al. (2002)). From this, we would
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50 484 expect that larger primates would have relatively larger and stronger forearm muscles;
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52 485 they would not, however, have relatively faster or more flexible muscles. This suggests
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486 that larger primates benefit more from stronger forearm muscles than they do from
487 faster-moving forearm muscles. With that said, in much of our previous work (e.g.
488 Hartstone-Rose et al. (2012a)), we have found positive allometry in muscle masses and
489 cross-sections, but isometry in fiber lengths – not necessarily because stretch/speed
490 simply scales directly with body size, but because FL tends to be where we see the
491 strongest functional signals. Thus, there is variation in FL between species because of
492 behavioral needs (e.g., frugivory vs. folivory adaptations in masticatory muscles), and
493 this variation reduces the coefficient of correlation sufficiently to reduce any positive or
494 negative allometry signal.

495 When considering the residuals data, there were no statistically significant
496 patterns independent of body mass supporting the existence of any taxonomic effects
497 within forearm muscle architecture.

498 Surprisingly, we also did not find architectural signals related to
499 locomotor/postural categorizations; forearm muscle fiber architecture does not reveal
500 statistical differences between quadrupedal primates and those that grip trees vertically
501 or leap propulsively. (Admittedly, that propulsion is achieved almost entirely by the
502 hind-limb musculature; see Marchi et al., this issue).

503 We did find a strong signal separating primates by substrate use: by some
504 metrics, terrestrial primates are significantly stronger than arboreal primates, and by
505 more substantial signals, arboreal primates have significantly greater FL than terrestrial
506 primates suggesting adaptations for speed and flexibility in the trees. These results
507 demonstrate how the functional tradeoff between strength vs. speed, a tradeoff that

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3 508 we have explored more extensively as we have related masticatory muscle fiber
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6 509 architecture to dietary specializations, applies to the forearm muscles. Although we
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9 510 could find no indication supporting our hypothesis that vertical clingers and leapers
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11 511 have relatively stronger muscles than those in other locomotor groups, it does seem
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13 512 clear that arboreality requires a different architecture than terrestriality.
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16 513 In the context of Jenkins' model (1973), it would be expected for groups such as
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18 514 arboreal, suspensory, and terrestrial primates to have larger digital flexor volumes and
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21 515 larger bony insertions because of their employment in locomotion. However, only our
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23 516 results for terrestrial primates fit this model, which suggests that the use of bony origins
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26 517 on the distal humerus may not always be an accurate method to predict the force
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28 518 production capabilities of the forearm musculature in primates.
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31 519 As an extension of this project, our lab is currently examining the covariation of
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33 520 muscle force production with the osteology of their origins – the epicondyles of the
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36 521 distal humerus. Future work will benefit from data on antebrachial muscle strength,
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38 522 allowing for reconstructions of the biomechanical loads resulting from muscle action
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41 523 around the joint, which may speak directly to the models proposed by Jenkins (1973).
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43 524 Although previous work has demonstrated that the length and retroflexion of the
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46 525 medial epicondyle may be a significant predictor of substrate use (arboreality v.
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48 526 terrestriality) in the fossil record, our findings imply a lack of support for hypotheses
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51 527 relating posture (i.e. quadrupedal/orthograde) and locomotion
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53 528 (quadrupedal/suspensory/vertical clinging and leaping) to the distal humerus, as we
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56 529 could not find the soft-tissue relationships that those hypotheses were based on.
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530 However, we are now collecting data on the humeri of the same individuals from this
531 study to test these hypotheses that have related that morphology to specific behaviors
532 requiring differences in strength.

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535

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542 full data set until the acceptance of our follow-up paper combining these soft-tissue
543 results with osteometrics, we are happy to freely share these data (along with data from
544 our other papers in this issue on leg and masticatory architecture of the same
545 specimens) upon request (please contact AHR); as these papers focus on functional
546 correlates of a broad section of primates with little examination of phylogenetic effects,
547 we are particularly interested in sharing our data with anyone interested in leading an
548 exploration of the phylogenetic effects of our forearm (perhaps in combination with the
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this study with their body mass, locomotor/postural category and substrate designations; species with

2 two individuals included in the sample are indicated by ⁺.

Species	Common Name	Superfamily ¹	Body Size (g) ²	Locomotion		Substrate
				Category ³	Subtype ⁴	
<i>Eulemur fulvus</i> ⁺	Brown Lemur	Lemuroidea	2215*	Quadrupedalism	Branch walking and running; Leaping; Vertical climbing; Suspension	Arboreal
<i>Eulemur macaco</i>	Black Lemur	Lemuroidea	1880	Quadrupedalism	Branch walking and running; Leaping; Vertical climbing; Suspension	Arboreal
<i>Haplemur griseus</i>	Gray Bamboo Lemur	Lemuroidea	709*	Vertical Clinging/Leaping	Leaping; Branch walking and running; Vertical climbing; Suspension	Arboreal
<i>Varecia rubra</i>	Red Ruffed Lemur	Lemuroidea	3300 ⁵	Quadrupedalism	Branch walking and running; Leaping; Vertical climbing; Feet suspension	Arboreal
<i>Microcebus murinus</i>	Gray Mouse Lemur	Lemuroidea	59	Quadrupedalism	Branch walking and running; Leaping	Arboreal
<i>Daubentonia madagascariensis</i>	Aye-Aye	Lemuroidea	2555.00	Tapper	Branch walking and running; Vertical climbing; Leaping	Arboreal
<i>Galago senegalensis</i>	Senegal	Lorisoidea	213*	Vertical	Leaping; Branch walking and running;	Arboreal

	Bushbaby			Clinging/Leaping		
<i>Nycticebus coucang</i>	Slow Loris	Lorisoidea	652.50*	Quadrupedalism	Quadrupedal walking; Vertical climbing	Arboreal
<i>Aotus nancymae</i>	Nancy Ma's Night Monkey	Ceboidea	787	Quadrupedalism	Quadrupedal walking; Leaping	Arboreal
<i>Aotus azarae</i> ⁺	Azara's Night Monkey	Ceboidea	1230	Quadrupedalism	Quadrupedal walking; Leaping	Arboreal
<i>Ateles fusciceps</i>	Black- Headed Spider Monkey	Ceboidea	8890.00	Suspensory ⁷	Brachiation; Leaping; Bipedal walking	Arboreal
<i>Callithrix jacchus</i> ⁺	Common Marmoset	Ceboidea	320.5	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Callithrix geoffroyi</i> ⁺	White- Headed Marmoset	Ceboidea	359	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Cebuella pygmaea</i> ⁺	Pygmy Marmoset	Ceboidea	116	Vertical Clinging/Leaping	Branch walking and running; Leaping; Vertical climbing	Arboreal

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<i>Leontopithecus chrysomelas</i>	Golden-Headed Lion Tamarin	Ceboidea	577.50*	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Leontopithecus rosalia</i>	Golden Lion Tamarin	Ceboidea	609*	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Saguinus bicolor</i>	Pied Tamarin	Ceboidea	429*	Quadrupedalism	Branch walking and running; Leaping	Arboreal
<i>Saguinus labiatus</i>	White-Lipped Tamarin	Ceboidea	539	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Saguinus oedipus</i>	Cotton-Top Tamarin	Ceboidea	404	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Saguinus imperator</i>	Emperor Tamarin	Ceboidea	475	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Saguinus midas</i>	Golden-Handed Tamarin	Ceboidea	515	Vertical Clinging/Leaping	Branch walking and running; Leaping	Arboreal
<i>Saimiri sciureus</i> ⁺	Squirrel Monkey	Ceboidea	723.50*	Quadrupedalism	Branch walking and running; Leaping	Arboreal

<i>Sapajus apella</i> ⁺	Tufted Capuchin	Ceboidea	3085*	Suspensory ⁷	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus hamlyni</i>	Hamlyn's Monkey	Cercopithecoidea	3360	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus cephus</i>	Moustached Guenon	Cercopithecoidea	4290	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus neglectus</i>	De Brazza's Monkey	Cercopithecoidea	10635	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus petaurista</i>	Lesser Spot-Nosed Monkey	Cercopithecoidea	2900	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus cambelli</i>	Campbell's Mona Monkey	Cercopithecoidea	1505	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercopithecus mona</i>	Mona Monkey	Cercopithecoidea	5100	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Chlorocebus aethiops</i> ⁺	Grivet	Cercopithecoidea	3620*	Quadrupedalism	Quadrupedal walking and running; Leaping	Terrestrial
<i>Erythrocebus patas</i>	Patas Monkey	Cercopithecoidea	8185*	Quadrupedalism	Ground walking and running (hand digitigrady)	Terrestrial

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<i>Miopithecus talapoin</i> ⁺	Angolan Talapoin	Cercopithecoidea	1380	Quadrupedalism	Quadrupedal walking and running; Leaping	Arboreal
<i>Cercocebus atys</i> ⁺	Sooty Mangabey	Cercopithecoidea	10350	Quadrupedalism	Ground walking and running (hand digitigrady)	Arboreal
<i>Cercocebus torquatus</i>	Collared Mangabey	Cercopithecoidea	6230	Quadrupedalism	Ground walking and running (hand digitigrady)	Arboreal
<i>Macaca sylvanus</i>	Barbary Maque	Cercopithecoidea	16000	Quadrupedalism	Quadrupedal walking and running	Arboreal
<i>Mandrillus sphinx</i>	Mandrill	Cercopithecoidea	6900	Quadrupedalism	Ground walking and running (hand digitigrady); Vertical climbing	Terrestrial
<i>Mandrillus leucophaeus</i>	Drill	Cercopithecoidea	20000	Quadrupedalism	Ground walking and running (hand digitigrady); Vertical climbing	Terrestrial
<i>Papio sp.</i>	Baboon	Cercopithecoidea	78200 ⁶	Quadrupedalism	Ground walking and running (hand digitigrady)	Terrestrial
<i>Colobus guereza</i>	Mantled Guereza	Cercopithecoidea	9200	Quadrupedalism	Quadrupedal walking and running; Bounding; Leaping	Arboreal
<i>Hylobates lar</i>	Lar Gibbon	Hominoidea	5340*	Suspensory ⁷	True Brachiation, Leaping, Bipedal walking	Arboreal
<i>Gorilla gorilla gorilla</i> ⁺	Gorilla	Hominoidea	120950 *	Suspensory ⁷	Ground walking and running (knuckle-walking); Vertical climbing; Modified Brachiation	Terrestrial
<i>Pan troglodytes</i>	Chimpanze	Hominoidea	45466*	Suspensory ⁷	Ground walking and running (knuckle-walking);	Terrestrial

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	e				Vertical climbing; Modified Brachiation	
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- ¹Ankel-Simons (2007).

²Fleagle (1999) except as noted; average mass for sex if sex known; average for species if sex unknown (annotated by “*”)

³Napier and Napier (1967) except as noted

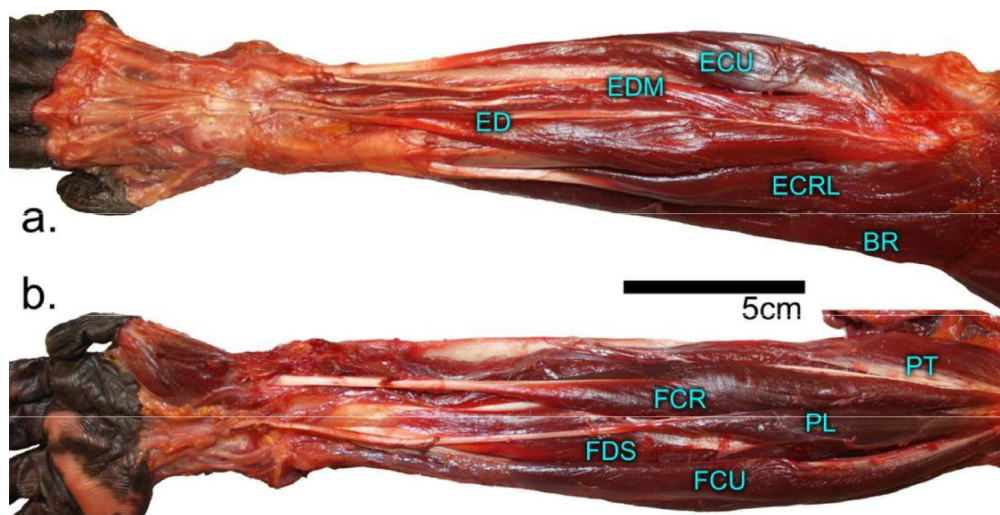
⁴Hunt et al. (1996)

⁵Body size for *Varecia variegata*

⁶*Papio* sp. body mass calculated as average mass for genus based on Fleagle (1999)

⁷Cant et al. (2003)

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Posterior (a) and anterior (b) views of a representative primate forearm, with the muscles of interest labeled.

55x28mm (600 x 600 DPI)

Table 2. Descriptive statistics for RMA regressions of architectural variables against body mass across the whole sample. As described in the methods, because the cube-root and square-root was taken of the volumetric and area variables respectively, all expected slopes = 1.

Y-variable	Slope (β) ^a	Y- intercept	r	Lower β CL	Upper β CL
Log Muscle Mass (g) ^{1/3}					
Total Flexors and					
Extensors	1.08	-0.79	0.98	1.02	1.16
Flexors	1.09	-0.85	0.98	1.02	1.17
Extensors	1.05	-0.91	0.97	0.97	1.13
Total Muscles	1.11	-0.78	0.98	1.04	1.18
Other Muscles	1.20	-1.14	0.96	1.09	1.32
Digital Flexors	1.07	-0.89	0.98	1.00	1.15
Wrist Flexors	1.18	-1.13	0.97	1.08	1.28
Digital Extensors	1.08	-1.08	0.96	0.98	1.19
Wrist Extensors	1.01	-0.96	0.96	0.92	1.12
Log FL (mm)					
Total Flexors and					
Extensors	0.93	0.22	0.96	0.84	1.02
Flexors	0.93	0.22	0.94	0.83	1.05
Extensors	0.98	0.16	0.90	0.83	1.15
Total Muscles	1.00	0.17	0.94	0.88	1.13

Other Muscles	1.30	-0.12	0.77	0.98	1.71
Digital Flexors	0.96	0.22	0.93	0.84	1.09
Wrist Flexors	0.86	0.21	0.88	0.72	1.03
Digital Extensors	0.90	0.19	0.90	0.77	1.06
Wrist Extensors	1.01	0.17	0.92	0.88	1.17

Log PCSA ^{1/2}

Total Flexors and					
Extensors	1.20	-0.83	0.97	1.10	1.31
Flexors	1.26	-0.99	0.95	1.14	1.40
Extensors	1.17	-1.04	0.95	1.05	1.30
Total Muscles	1.24	-0.82	0.97	1.14	1.35
Other Muscles	1.33	-1.27	0.94	1.17	1.50
Digital Flexors	1.18	-1.01	0.95	1.06	1.32
Wrist Flexors	1.47	-1.46	0.94	1.29	1.66
Digital Extensors	1.33	-1.40	0.92	1.15	1.54
Wrist Extensors	1.13	-1.16	0.92	0.98	1.29

Log RPCSA ^{1/2}

Total Flexors and					
Extensors	1.24	-0.89	0.97	1.14	1.35
Flexors	1.30	-1.05	0.97	1.19	1.43
Extensors	1.15	-1.03	0.96	1.04	1.27
Total Muscles	1.25	-0.87	0.97	1.15	1.36
Other Muscles	1.31	-1.38	0.94	1.15	1.50
Digital Flexors	1.28	-1.16	0.93	1.12	1.47
Wrist Flexors	1.46	-1.43	0.96	1.31	1.62

Digital Extensors	1.20	-1.27	0.93	1.05	1.39
Wrist Extensors	1.13	-1.14	0.95	1.00	1.27

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Figure 2. shows the Flexor FL plotted against Log BM for arboreal and terrestrial primates. The black line indicates the orthogonal fit across the whole sample; the blue line indicates fit for arboreal species and the red line indicates fit for terrestrial species. Strepsirrhines are denoted as green shapes, Platyrrhines as blue shapes, and Catarrhines as red shapes. Open shapes indicate terrestrial species; filled shapes indicate arboreal species. Quadrupedal primates are denoted by squares, VCL primates are denoted by triangles, and suspensory primates are denoted by circles. *Daubentonia madagascarensis* alone is indicated by an asterisk, as it is classified as a tapper.

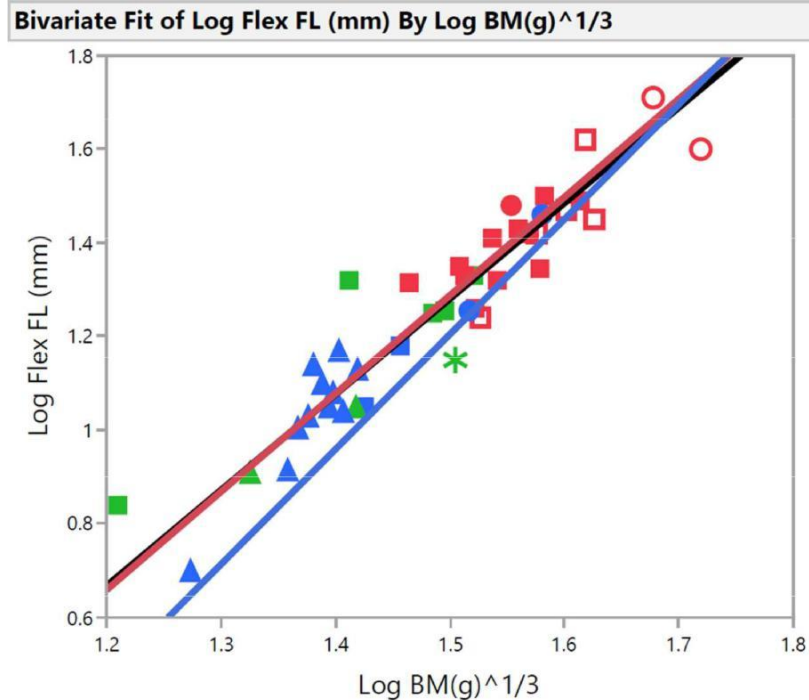


Table 3. Descriptive statistics for one-way analyses of residuals of architectural variables grouped by substrate use. P-values annotated by “*” are significant at an alpha of 0.05, “**” alpha >0.01 and those annotated “~” approach significance.

	Substrate Use		
	Means		p-value
	Arboreal	Terrestrial	
	Log Muscle Mass(g)^1/3		
Total Flexors and Extensors	0.00	0.01	0.40
Flexors	0.00	0.01	0.40
Extensors	0.00	0.00	0.84
Total Muscles	0.00	0.01	0.48
Other Muscles	0.00	-0.01	0.63
Digital Flexors	0.00	0.01	0.40
Wrist Flexors	0.00	0.01	0.51
Digital Extensors	0.00	0.00	0.96
Wrist Extensors	0.00	0.00	0.98
	Log FL(mm)		
Total Flexors and Extensors	0.00	-0.03	0.01*
Flexors	0.00	-0.02	0.11
Extensors	0.00	-0.02	0.25
Total Muscles	0.01	-0.04	0.01*
Other Muscles	0.02	-0.10	0.009**
Digital Flexors	0.01	-0.04	0.04*

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Wrist Flexors	0.00	-0.03	0.14
Digital Extensors	0.01	-0.03	0.12
Wrist Extensors	0.01	-0.03	0.13

Log PCSA^{1/2}

Total Flexors and Extensors	0.00	0.02	0.10~
Flexors	0.00	0.02	0.34
Extensors	0.00	0.00	0.85
Total Muscles	0.00	0.02	0.09~
Other Muscles	0.00	0.00	0.88
Digital Flexors	0.00	0.02	0.20
Wrist Flexors	0.00	0.01	0.75
Digital Extensors	0.00	-0.01	0.72
Wrist Extensors	0.00	0.00	0.98

Log RPCSA^{1/2}

Total Flexors and Extensors	0.00	0.03	0.03*
Flexors	0.00	0.04	0.05*
Extensors	0.00	0.02	0.20
Total Muscles	0.00	0.03	0.09
Other Muscles	0.00	0.02	0.48
Digital Flexors	0.00	0.03	0.26
Wrist Flexors	0.00	0.01	0.59
Digital Extensors	0.00	-0.02	0.43
Wrist Extensors	0.00	0.04	0.08~