

1 **A novel biomechanical model of the mouse forelimb predicts muscle activity in optimal control**
2 **simulations of reaching movements**

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15 **ABSTRACT**

16 Mice are key model organisms in genetics, neuroscience and motor systems physiology. Fine motor control
17 tasks performed by mice have become widely used in assaying neural and biophysical motor system
18 mechanisms, including lever or joystick manipulation, and reach-to-grasp tasks (Becker et al., 2019; Bollu
19 et al., 2019; Conner et al., 2021). Although fine motor tasks provide useful insights into behaviors which
20 require complex multi-joint motor control, there is no previously developed physiological biomechanical
21 model of the adult mouse forelimb available for estimating kinematics (including joint angles, joint
22 velocities, fiber lengths and fiber velocities) nor muscle activity or kinetics (including forces and moments)
23 during these behaviors. Here we have developed a musculoskeletal model based on high-resolution imaging
24 and reconstruction of the mouse forelimb that includes muscles spanning the neck, trunk, shoulder, and
25 limbs using anatomical data. Physics-based optimal control simulations of the forelimb model were used
26 to estimate *in vivo* muscle activity present when constrained to the tracked kinematics during mouse
27 reaching movements. The activity of a subset of muscles was recorded via electromyography and used as
28 the ground truth to assess the accuracy of the muscle patterning in simulation. We found that the synthesized
29 muscle patterning in the forelimb model had a strong resemblance to empirical muscle patterning,
30 suggesting that our model has utility in providing a realistic set of estimated muscle excitations over time
31 when provided with a kinematic template. The strength of the resemblance between empirical muscle
32 activity and optimal control predictions increases as mice performance improves throughout learning of the
33 reaching task. Our computational tools are available as open-source in the OpenSim physics and modeling
34 platform (Seth et al., 2018). Our model can enhance research into limb control across broad research topics
35 and can inform analyses of motor learning, muscle synergies, neural patterning, and behavioral research
36 that would otherwise be inaccessible.

37 INTRODUCTION

38 Mice are ubiquitous model organisms across many fields of biological research, including genetics,
39 neuroscience and physiology. They provide access to a wide array of disease models, advanced molecular
40 interrogation techniques, ease of husbandry through short gestation cycles, and the cost of raising and
41 housing mice is relatively inexpensive. In addition to these factors, mice are often used in behavioral
42 studies, including those that assay fine motor control. Mice can perform tasks like manipulandum control
43 (Bollu et al., 2019), dexterous reach (Becker et al., 2020), and can learn complex behaviors with training
44 (Burgess et al. 2017, Serradj et al., 2023, Sauerbrei et al., 2020, Galinanes et al., 2018, Conner et al., 2021).
45 However, despite the utility of mice as a model organism in motor learning, there are no high-resolution
46 reconstructions of the adult mouse forelimb, nor are there any physiological biomechanical models of their
47 forelimbs that incorporate fully developed muscle morphology. Biomechanical models are useful for motor
48 systems and neuromechanics researchers to provide detailed insights into muscle activity and limb
49 kinematics (e.g., fiber length and velocities) that would otherwise be difficult or impossible to access
50 through empirical observations. Experimentally measuring muscle activity is challenging due to the size of
51 the mouse and the large number of muscles in the body. State-of-the-art methods can only measure the
52 activity of 3-4 muscles in the 25+ muscles in the forelimb (Zia et al., 2020). Therefore, the construction and
53 evaluation of a model of the mouse forelimb would be a valuable tool for researchers studying dexterous
54 behaviors in mice.

55 The only currently available mouse forelimb model, developed recently in a full-body mouse model
56 (Ramalingasetty et al., 21), was based on mouse embryo data (Delaurier et al., 2008), which also lacked
57 many of the large muscles originating from the scapula, and on educated guesses. Ramalingasetty et al.
58 noted that modeling the mouse forelimb is more challenging than the hindlimb, due to the lack of published
59 biomechanical data, and improving their forelimb model was identified as a remaining challenge for future
60 work. Reference books on limb anatomy present two-dimensional (2D) illustrations of the limb musculature
61 (Hebel 1986), but it is challenging to extract accurate locations of the attachment points and the three-
62 dimensional (3D) tissue paths from these references (Delaurier et al., 2008). By using large-scale light sheet
63 microscopy data, we were able to more accurately identify the muscle attachment sites and muscle paths
64 than by working with mouse and rat atlas data. Additionally, computing these quantities directly from
65 dissections is challenging because of the size of the forelimb muscles, whose tendon insertion points are
66 separated by as little as tens of microns. The attachment points have been shown to be the most important
67 factor in estimating how effective a muscle is in producing a joint rotation or moment (Charles et al., 2016).

68 The study by Charles et al. has produced a detailed description of muscle anatomy integrated into
69 a hindlimb biomechanical model. We sought to build on this work to create a forelimb model. We started
70 by scanning and recreating the forelimbs of two adult mice. Muscles with insertions onto the humerus span
71 most of the mouse's trunk and spine, necessitating imaging of much of the mouse body. We limited our
72 reconstruction to muscles that had insertions onto the humerus, radius, and ulna, as reconstruction of
73 muscles with insertions onto the scapula and those that inserted onto the hand was infeasible given the
74 resolution of imaging performed. Once the muscles had been traced and reconstructed, they were used to
75 set the musculoskeletal geometry of the biomechanical model, that is, the attachments points of the muscles
76 on the bones and their lines of action. We used published results on mouse forelimb muscle architecture to
77 set the muscle parameters in our model (Mathewson et al., 2012). The resulting model has 21 muscles and
78 5 bones (along with a composite hand body segment), with the scapula and clavicle serving as fixed position
79 bodies. The model has four degrees-of-freedom: shoulder elevation, extension, and rotation, as well as
80 elbow flexion. The model is also capable of wrist flexion and rotation, but these degrees-of-freedom were
81 fixed during our simulations. We used the OpenSim modeling and physics simulation environment to
82 develop the forelimb model (Delp et al. 2007, Kewley et al. 2024). We have also written custom code to
83 convert the model for the MuJoCo physics simulation environment (Todorov et al. 2012).

84 To evaluate the utility of the model, we sought to replicate physiological kinematics and predict
85 simultaneously recorded muscle activity. We used a dataset of thousands of reaches from three mice who
86 had their kinematics and a subset of their muscle activity recorded during reaching movements. The
87 empirical kinematics were used as constraints on the synthesized kinematics with optimal control. The

88 empirical muscle activity was used as a ground truth for comparison against the optimal control predictions
89 (i.e., the synthesized muscle activity). The muscles recorded experimentally in the dataset were the biceps,
90 triceps long head and triceps lateral head.

91 Reaching movements in mice, rats, and primates are widely studied in systems neuroscience to
92 study healthy and injured motor networks (Fleischer et al., 2023; Khanna et al., 2021; Yang et al., 2023).
93 However, reaching movements lack endpoint accuracy in patients with cerebellar disease (Bonnetfoi-
94 Kyriacou et al., 2018), exhibit abnormal muscle coordination patterns after stroke (Cheung et al., 2012) and
95 have impaired kinematics in Parkinson's disease (Vissani et al., 2021). The mouse is an ideal model system
96 for motor control research because of relatively easy access to neural, behavior and anatomical data, as well
97 as to advanced molecular interrogation techniques for perturbation studies (Deisseroth et al., 2006).
98 Building on a large body of prior work in primates, several research groups are conducting foundational
99 studies on mouse reaching (Becker et al., 2020; Yang et al., 2022; Wagner et al., 2021; Conner et al., 2021;
100 Galilanes et al., 2018), which are evolutionarily conserved (Iwaniuk 2000). Reaching movements are the
101 focus of the model evaluation experiments in this study; however, the model could be used to simulate other
102 forelimb movements.

103 There is an infinite number of possible muscle coordination patterns that are consistent with
104 kinematics (Harris & Wolpert 1998). Optimal control chooses the muscle excitation pattern that achieves
105 task constraints, while minimizing a proxy for effort or energy (e.g., the sum of muscle activations squared)
106 and possibly other terms (Al Borno et al., 2020). We apply optimal control to predict an energetically
107 efficient muscle activity pattern that achieves the reaching kinematics task. We focused this study on the
108 ballistic phase of the reach and have not studied the grasping phase. To simplify the problem, we have not
109 included the muscles that control the wrist and fingers in the biomechanical model and kept these degrees-
110 of-freedom locked. However, we have provided the 3D reconstruction of some of these muscles in
111 supplementary materials. We are not aware of any prior work that compared predicted muscle activity with
112 empirical muscle activity for three-dimensional reaching movements (in humans or other species).

113 Optimal control-based simulations using the model were able to recreate reach kinematics
114 accurately using synthesized muscle excitations. The muscle patterning produced when constrained to
115 replicate experimental reach kinematics had a strong resemblance to empirical electromyography (EMG)
116 data. The model performed best when estimating the mean EMG rather than on a reach-per-reach basis
117 because of the high physiological variance in the muscle patterning employed in mice. Mean EMG
118 predictions were within 1 standard deviation of mean experimental EMG in most reaches and produced
119 lower error than time-shuffled EMG. These results suggest that our model can replicate realistic reach
120 kinematics and muscle activity. Our analysis reveals that the optimal control solutions are closer to the
121 empirical solutions (i.e., the patterns employed by real mice) as reaching performance improves throughout
122 learning. In other words, mice employ muscle patterning solutions that more closely resemble optimal
123 control solutions as they become more skilled at the task. More broadly, this model should provide insight
124 into forelimb behaviors that would otherwise be inaccessible by experimental means, and we hope that
125 access to a robust description of the forelimb's kinematics, forces, and muscle activity will advance
126 understanding of mouse behavior. Our computational tools are available as open-source for researchers
127 interested in analyzing muscle activity during mouse forelimb movements.

128

129 **METHODS**

130 **Anatomical high-resolution imaging**

131 Accurate prediction of muscle activity during movements is predicated on a sufficient description of the
132 underlying anatomy and physiology. To produce a model that was suited for prediction tasks, we first sought
133 to gather anatomical data to inform the model. We obtained 3D scans of mouse forelimbs and trunk
134 musculature through large scale light sheet microscopy imaging of two wildtype female mice (9 weeks old).
135 The dataset contains imaging that captured the left distal shoulder and proximal forelimb (Mouse A), the
136 right distal forelimb and paw (Mouse A), and both forelimbs, shoulders, and trunk (Mouse B). Only the left
137 shoulder, trunk, and proximal forelimb were reconstructed in Mouse B.

138

139 **Mouse and tissue preparation**

140 Mice were euthanized via subcutaneous injection of pentobarbital and prepared for imaging through fixation
141 with a transcardial injection of 4% paraformaldehyde (PFA) and 0.01% heparin. The vascular system was
142 washed with a saline solution before and after perfusion with PFA, then washed overnight in phosphate-
143 buffered saline (PBS) and 0.01 heparin. Mouse A was dissected, with skin removed and forearm separated
144 axially to the vertebral column.

145 Imaging subjects were prepared using the iDISCO+ tissue clearing methodology (Habart et al.,
146 2023). The tissue was introduced to a gradually increasing concentration of methanol, starting with 20%
147 and increasing by 20% every hour. The clearing chamber was maintained at room temperature. After 5
148 hours of exposure to methanol, the tissue was chilled at 4° C overnight and then bathed in 66%
149 dichloromethane (DCM) and 33% methanol for 24 hours. The tissue was then bathed in 100% methanol
150 for 2 hours before being chilled for 1 hour and then transferred into 5% hydrogen peroxide in methanol for
151 48 hours. Finally, tissue was rehydrated through 1 hour immersion in 80%/60%/40%/20% methanol for one
152 hour per 20% decrement, then transferred to 1x PBS for 24 hours, followed by immersion in a 100ml PBS
153 10x and 2 ml TritonX-100 solution that was filled to 1L with distillate water.

154 After clearing was completed, the tissue was prepared for immunostaining without antibodies via
155 incubation a permeabilization solution (500 mL) consisting of 400 mL PTx.2, 11.5 g glycine, and 100 mL
156 dimethylsulfoxide (DMSO). The tissue was bathed in solution for 4 days, then transferred to a blocking
157 solution of 42 mL PTx.2, 3 mL donkey serum, and 5 mL DMSO for 3 days. Finally, tissue was washed with
158 100 mL PBS 10X, 2 mL Tween-20, 1 mL of 10mg/mL heparin, and filled to 1L with distillate water. The
159 tissue was then re-cleared through preparation in 20%/40%/60%/80%/100% methanol in 1-hour steps, then
160 bathed in 100% methanol overnight. Afterwards, the tissue was bathed in 66% DCM and 33% methanol
161 for 4 hours, then in 100% DCM for 15 minutes twice in succession.

162

163 **Imaging parameters**

164 Dissected mouse limbs were arranged in a prone position prior to imaging. Scans were taken with 8.23 um
165 per pixel scans at 8x zoom, with 5 um steps in the z-plane. Imaging was performed using mesoSPIM (Voigt,
166 et al., 2019). Immunostaining was captured in the green channel (488 nm laser) and was imaged using mode
167 tiling wizard with an offset by 75% and a filter set to 530/43. Mouse A's forearms were dissected and imaged
168 in their entirety. Mouse B was imaged from the base of the skull through the joint of the femur and tibia
169 and the entirety of the depth of the sample.

170

171 **Anatomical segmentation and reconstruction**

172 The basis for this study was to obtain physiological morphology data from the mouse forelimb to inform
173 the development of a musculoskeletal model for usage in optimal control-based simulations. We found that
174 muscle density and striation was a sufficient marker of muscles to identify them with light sheet microscopy,
175 which were enhanced through immunostaining without antibodies (see **Methods: Mouse and tissue**
176 **preparation**). We used the raw imaging of the mouse anatomy and segmented individual muscles into 3D
177 shape objects using 3D Slicer (Federov et al., 2012) (Fig. 1A). We also segmented the forelimb bones to
178 obtain landmarks and geometries for use in the model, such as the deltoid tuberosity of the humerus, which
179 is a site of attachment for many shoulder muscles in the mouse. Because not every data set had complete

180 data for the entire forelimb, right and left anatomical datasets were combined through manual alignment in
181 Blender (Blender D.T., 2022). We used anatomical landmarks on the humerus, ulna, and radius to align
182 muscle reconstructions, as these bones were present in all three imaging datasets. The reconstructions
183 were scaled according to the radius of the bones and confirmed visually by examining the degree of overlap
184 between reconstructions.

185

186 **Development of a biomechanical model of the mouse forelimb**

187 With a detailed set of reconstructions, we next sought to leverage anatomical descriptions to construct a
188 biomechanical model in OpenSim, a widely-used physics-based modeling and simulation environment used
189 to study movements of humans and other species. The anatomical model was assembled using OpenSim
190 Creator (Kewley et al., 2024). We have also converted the model in MuJoCo (Todorov et al. 2012), which
191 produces faster (but less physiologically accurate) simulations that are more amenable for deep
192 reinforcement learning applications. Although the model is available in MuJoCo, the computational tools
193 for optimal control are based in OpenSim; therefore, MuJoCo users will need to develop their own code to
194 produce the simulations with the model. Each individual muscle was measured and a combination of
195 parameters derived from optical measurement, and from previous parameters in Mathewson et al., 2012
196 and Charles et al., 2016, were used to derive the biophysical properties of the modelled muscles. We used
197 De Groote-Fregly (De Groote et al., 2016) Hill-type muscles within the model, and opted to ignore complex
198 tendon dynamics (i.e., using rigid tendons with no force-length/velocity properties), both to facilitate the
199 production of a functional model and because we did not have access to sufficient data regarding tendon
200 physiology purely from imaging data. The model is likely to improve from a more detailed dissection and
201 biophysical tests, but these assays were beyond the scope of this project.

202 The musculotendon units in the forelimb were modeled by the common Hill-type muscle (Uchida
203 & Delp 2021), which is parametrized by four parameters (maximum isometric force, optimal fiber length,
204 tendon slack length and pennation angle). All these parameters, except for tendon slack length, were
205 determined experimentally in the muscle dissection study of Mathewson et al., 2012 and through
206 interpolation from known values when a muscle was not described in prior literature. The tendon slack
207 length parameter represents the length where a tendon develops passive elastic force (Uchida & Delp 2021).
208 This parameter cannot be measured experimentally and was set using the optimization procedure of
209 Buchanan et al., 2004, as is commonly performed in the field (Charles et al., 2016), assuming that muscle
210 fibers remain within 0.5 to 1.5 times optimal fiber length throughout the joint's range of motion, which
211 were estimated from both anatomical constraints and video of mouse behavior. Based on the muscle paths
212 from the digital segmentation, we used wrapping surfaces, which are geometric objects in OpenSim, to
213 constrain the muscles to have realistic paths of action. This is necessary for the model to produce realistic
214 moment arms (Charles et al., 2016). We set other parameters in the muscles such as the maximum
215 contraction velocity, the activation time constants and the force-length curves scaled based on prior work
216 on mouse physiology (Charles et al., 2016; see open-source model for details). We calculated the
217 physiological cross-sectional area (PCSA) by the standard formula developed by Alexander & Vernon
218 (1975), that is, muscle volume divided by fiber length. Muscle fiber pennation angle is entered separately
219 in OpenSim models; thus not directly used in PCSA calculations. Bone volume was determined in
220 reconstruction and was uniformly multiplied by a murine bone density scalar (.00425 kg/cm²) determined
221 from a literature search for empirical measures (Robbins et al. 2018) and prior models of the mouse (Charles
222 et al., 2016), as well as estimations of the center-of-mass and inertia. A description of the model geometry
223 is available in Table 1 and the muscle parameterization in Table 2.

224

225 **Model scaling**

226 Individual mice have variable limb dimensions that models must be altered to accommodate. We
227 accomplished this by using the scale tool in OpenSim to automatically scale the mass, length, and muscle
228 parameters of the model to fit the observed kinematic data originating from a particular mouse subject. We
229 used DeepLabCut (Mathis et al., 2018) to estimate paw, elbow and shoulder markers from video. Our scripts
230 adjusted the marker positions based on a 2D skeletal model with estimated limb lengths (derived from mean

231 inter-marker distances). These adjusted marker positions were then used to scale the OpenSim model to the
232 mouse's proportions.

233

234 **Mouse behavior**

235 Evaluation of the model against a behavioral dataset with a known ground truth, both in terms of kinematics
236 and muscle activity, is necessary to assay its utility. Mouse behavior was recorded from two cameras during
237 reaching behaviors and then processed using the DeepLabCut 3D motion tracking software (Mathis et al.,
238 2018). We collected data in a forelimb reaching task and recorded electromyography (EMG) from the biceps
239 brachii, triceps long head, and triceps lateral head. The activity of three muscles were measured
240 simultaneously with Myomatrix arrays (Zia et al., 2020). We estimated the elbow joint angle from the 3D
241 markers. We used the average limb lengths to adjust the DeepLabCut paw, elbow and shoulder markers and
242 ensure that the limb lengths remain constant throughout the video, which is necessary for accurate tracking
243 by the model. Because our forelimb model only has rotational degrees-of-freedom on the shoulder, we
244 could not capture the small translational movement occurring at the shoulder during head-fixed reaching.
245 We subtracted the shoulder markers displacements from the elbow and paw markers to keep shoulder
246 positional coordinates fixed in our simulations.

247 Processed EMG envelopes were normalized to the maximum contraction recording during the
248 session. EMG is usually normalized to the maximum voluntary contractions in studies with human subjects
249 ([Kendall et al., 2005](#)). We rectified the EMG signals and then filtered them with a bandpass and lowpass
250 filter suite. We bandpassed the signal from 5 to 500 Hz, rectified the signal, then low-passed further with a
251 cutoff of 10 Hz. Additionally, we normalized the filtered EMG signals with a z-score measure. Each muscle
252 was recorded through 4 leads, but only the qualitatively determined cleanest lead per muscle was used for
253 this study.

254 Mice were kept at 80% body weight during their training and testing periods, and mice were head-
255 fixed to a behavioral platform while reaching for small pellets (Figs. 1D, 2A). Mouse EMG was recorded
256 from session one, when the mouse was completely naïve, and training progressed indefinitely until expertise
257 was reached. The mice in this dataset ranged from having 11 to 26 total sessions, up to 1 hour per session.
258 The mice used for this study achieved an initial successful reach on a range of sessions spanning 2-5 days.

259

260 **Selection of reaches for simulation**

261 Our dataset spanned the entirety of reach training for 3 mice, and because of the progression of learning,
262 there was natural variance in kinematics performed. We opted to select only from 'expert' mice and to use
263 baseline EMG datasets that were derived from similar reaching kinematics. We grouped reaches using the
264 2-norm metric on 3D paw kinematics to assess similarity, and then selected two sets of 10 reaches per mice,
265 with each set having a different kinematic profile (i.e., qualitatively different paw trajectories). We enforced
266 expertise by selecting reaches that occurred only after the initial 4 sessions of learning, which was a typical
267 epoch for mice to reach moderate success in reaching. We also compared the optimal control predictions
268 between the early and late sessions of learning. Early sessions were selected from the 3 mice discussed
269 above, with an additional mouse who did not achieve expert status included. 10 reaches were selected from
270 each mouse for the early dataset. Early sessions were restricted to the first 3 sessions of learned reaching.

271

272 **Optimal control**

273 To demonstrate the ability of the model to synthesize physiological movements, kinematics were recorded
274 using DeepLabCut (Mathis et al., 2018). These provided 3D coordinates of the paw and elbow during head-
275 fixed reaching movements. Optimization was conducted with direct collocation in Moco (Dembia et al.,
276 2020) as it is well-suited for simulations that track experimental data ("inverse simulations"; e.g., Bishop
277 et al., 2021). Direct collocation enforces the equations-of-motion and physiological relationships as
278 constraints in a nonlinear optimization problem which solves for the states of the musculoskeletal system
279 and the muscle activity over the duration of the simulation. The optimization's objective is to minimize a
280 cost function of two terms: one term that is a proxy for effort (i.e., the sum of muscle activations squared)

281 and one term that represents the tracking cost (i.e., the deviation between the synthesized and the
282 experimental kinematics). The cost function equation is:

283

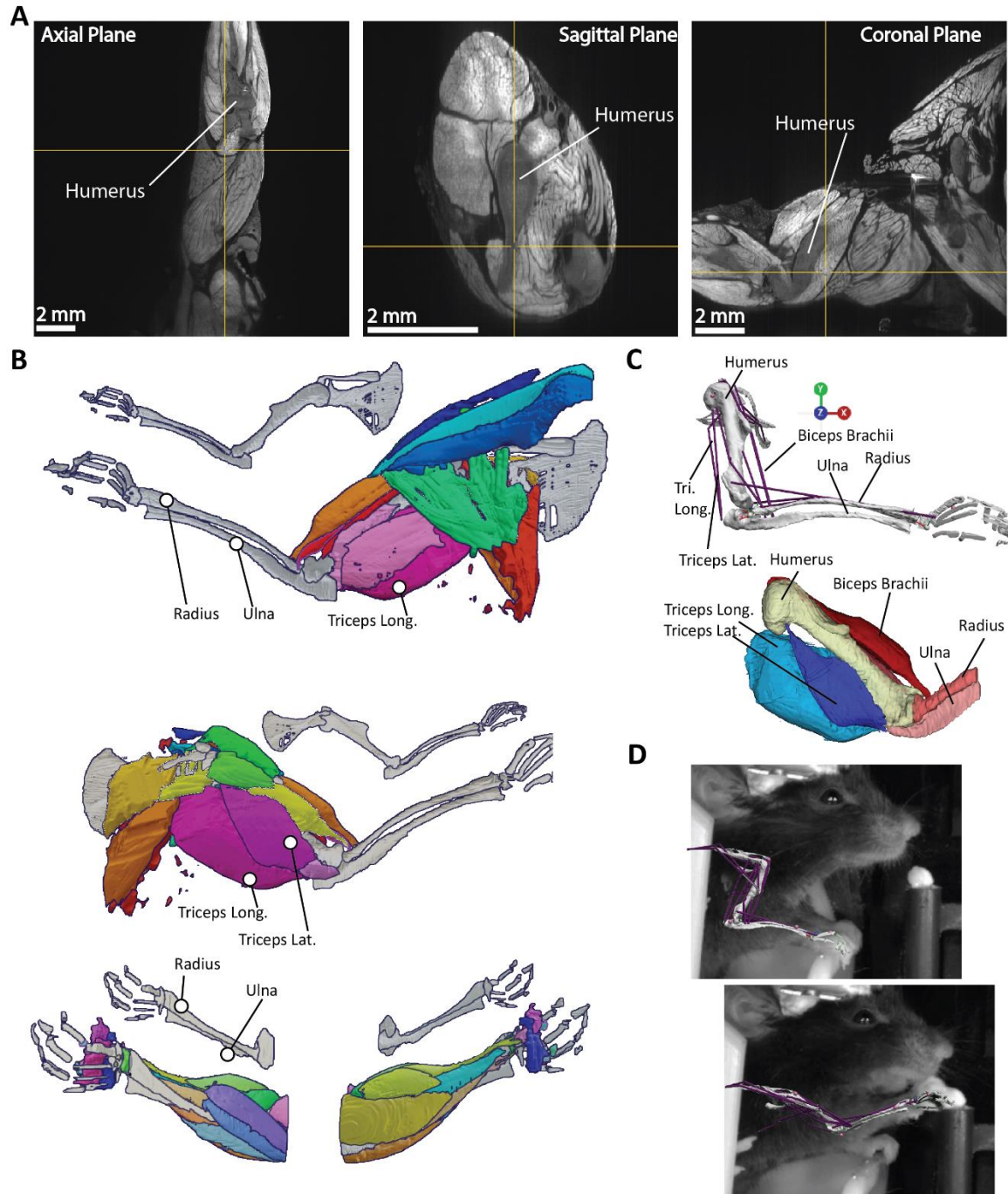
$$284 \quad E = wT_{\text{paw}} + wT_{\text{elbow}} + \sum_i a_i^2$$

285

286 Here, T_{paw} is the 2-norm squared difference between model and experimental 3D paw coordinates.
287 The experimental coordinates comprised of 100 timepoints during the ballistic phase of reaching. The same
288 holds for T_{elbow} , which is derived from tracking of the mouse elbow position. Term a_i denotes the activation
289 of muscle i in the model and w is a scalar weight set to 10^9 . We optimized over 2500 iterations and 100
290 mesh points. The simulation was also constrained to start and end with the joint angles derived during the
291 scaling of model. The optimization would end early if a convergence tolerance of $1e^{-7}$ was reached. The
292 optimization typically ran for 10 minutes on a computer with specifications listed in supplemental Table 1.

293

294 The empirical muscle activity was not fitted or used by the optimization. Muscle activity is
295 predicted based on optimality, task, physics and physiological constraints. We compared the predicted
296 muscle activity with electromyography measurements on the triceps lateral head, triceps long head and
297 biceps in 3 mice. We compared the activity of the recorded muscles to the muscle excitations produced by
298 the model with the mean absolute error (MAE) metric at an optimal lag (in a range of -50 to $+10$ ms; we
299 used a lag of 0 ms for Figure 3 and the late reach set in Figure 4; early reaches had an optimal lag of -50
300 ms in Figure 4). As a validation of the optimal control solutions, we compare the synthesized kinematics
301 between the musculoskeletal model and a torque-based model without muscles, but with motors on the
302 joints. We verified that the kinematics in muscle-based solutions closely recapitulated those in torque-based
303 solutions, indicating that the optimization converged to adequate kinematic solutions (i.e. kinematic means
304 were within a single standard deviation of the true kinematic mean across all dimensions, Figure 2: blue
305 violin plots).



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Figure 1. Anatomical reconstruction. **A.** Optical slices of the mouse forelimb in the axial, sagittal, and coronal planes. The mouse arm is oriented in the prone position. Labels added to highlight prominent muscles as an example of a reconstruction target. **B.** 3D projections of optical tracing results as a composite across mice. Upper panels show composite scan, while lower panels show the left hand of mouse A to highlight density of wrist-inserting muscles. 3D projections show morphology and attachment sites of muscles on bones that were used to create biophysical model. **C.** Biomechanical model (OpenSim) reconstruction developed from the 3D projections. **D.** Biomechanical model projection on video of mouse reaching.

315 RESULTS

316 Mouse forelimb musculoskeletal anatomy and biomechanical model construction

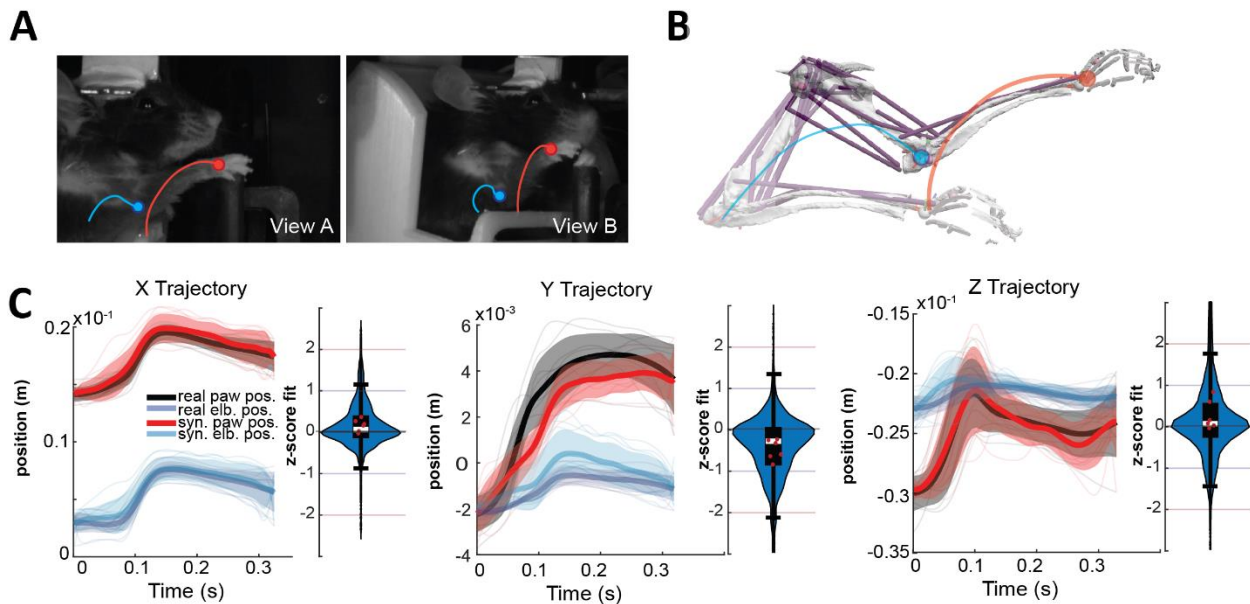
317 Table 1 and 2 provide data on the muscles segmented in this study, including their origins, insertions, and
318 physiological properties which were derived from optical scans (Fig. 1A-C). Overall, we segmented over
319 21 muscles and 5 bones. The constructed biomechanical model has 21 muscles, 4 degrees-of-freedom (Fig.
320 1C) and was sufficient to describe ballistic reaching movements (Fig. 1D).

321 Kinematic Tracking

322 We tasked the physiological forelimb model to track recorded DeepLabCut-tracked kinematics using
323 optimal control algorithms (see **Methods: Optimal control**). Because there was natural variance in
324 reaching movements and motor control, we opted to group 6 sets of 10 reaches by their kinematic
325 similarity across time (see **Methods: Selection of reaches for simulation**). We deliberately selected 6 sets
326 of varying reaching kinematics to explore the ability of the model to predict varied motor behaviors. The
327 model was scaled and then optimized with direct collocation (using OpenSim Moco, Dembia et al., 2021)
328 to track the paw and elbow across the ballistic epoch of the reach (Figure 2). We were able to recreate
329 limb kinematics with low error, with the majority of synthesized kinematics per timestep falling within 1
330 standard deviation of the empirical kinematic mean (Fig. 2C, blue violin plots; N = 60 reaches) across the
331 x, y, and z dimensions of the paw and elbow trajectories.
332

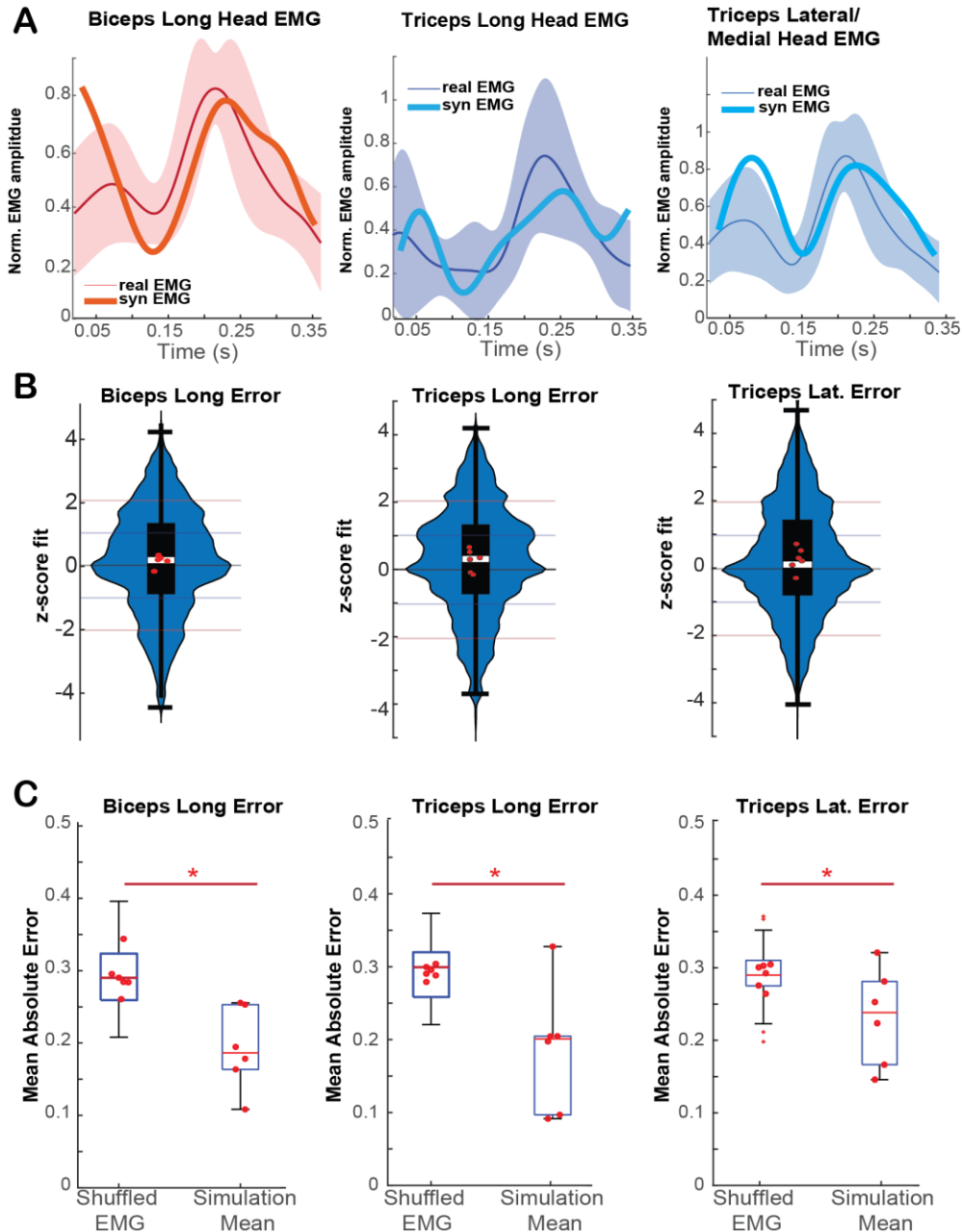
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337 **Figure 2. A.** Example video with schematic elbow and paw marker trajectories. **B.** A biomechanical model with virtual
338 markers on the elbow and paw. An optimal control problem is solved to minimize the difference between the virtual
339 and empirical markers. **C.** Mean and standard deviation of 60 reaches for the empirical and synthesized marker
340 trajectories. The z-score of the synthesized markers are largely within 1 standard deviation (see violin plots in blue),
341 and the means per set of 10 reaches are all within 1 standard deviation (red dots). Black box plots denote median
342 (white bar), 25 to 75th percentile distributions (black box), and 10th to 90th percentile distributions (short horizontal
343 black lines).



343

344 **Figure 3. A.** Comparisons of synthesized muscle excitations and experimental EMG activity. Curves show means and
345 standard deviations (line and shaded region) of 10 reaches with similar experimental elbow and paw trajectories that
346 were chosen from mouse behavior dataset. The mean synthesized excitations are shown in thick red for biceps long
347 head, thick cyan for triceps long head, and thick blue for triceps medial head compared to the base lateral head
348 activity. **B.** Violin plots including the entire dataset of 60 reaches. The mean synthesized muscle activity lies largely
349 within 1 standard deviation of mean experimental muscle activity (red dots). On a reach-by-reach basis, the
350 synthesized muscle activity lies largely within a z-score of 2 standard deviations (blue violin plots). Black box plots
351 denote median z-deviation (white bar), 25 to 75th percentile distributions (black box), and 10th to 90th percentile
352 distributions (horizontal black line). **C.** A comparison of mean absolute error between time-shuffled physiological
353 EMG data and synthetic excitation means to the real mean of the tracked data. Synthetic excitation means have lower
354 MAE than the time-shuffled data in all three muscles recorded (two-sided t-test, biceps $p = 2.2e-6$, triceps long head
355 $p = 1.4e-6$, triceps lateral head $p = 4.8e-4$. P-values were Holm-Bonferroni corrected for multiple comparisons). Black

356 box plots denote median (red horizontal bar), 25 to 75th percentile distributions (black box), and 10th to 90th
357 percentile distributions (short horizontal black lines and stems).

358 **Model muscle activity patterning for reaching movements**

359 We tasked the model to synthesize muscle activity during reaching movements with optimal control. We
360 experimentally recorded activity from the biceps, triceps lateral head, and triceps long head in 3 mice. We
361 then synthesized muscle activity to recapitulate experimental kinematics with direct collocation using
362 OpenSim Moco. As shown in the examples in Fig. 3A, we observe that the mean synthesized muscle activity
363 closely resembles empirical muscle activity over the duration of the reach for all three muscles. We measure
364 the performance of the model via the MAE of normalized ground truth EMG signals from model signals.
365 The mean muscle excitations produced by the model were within a single standard deviation of the
366 experimental EMG activity (Fig. 3B, red dots in violin plots; between 50-57 of the 60 reaches,
367 depending on the mice and assayed muscle). On a reach-
368 by-reach basis, the muscle excitations produced by the model
369 across all time points were typically within two standard
370 deviations of the experimental EMG activity. Paired reach-to-
371 reach predictions were less accurate because mice were highly
372 variable in their muscle patterning for the same kinematic
373 profile (and some muscle patterns may be inefficient, more
374 consistent with early learning or motor exploration, which
375 would not be predicted as accurately by optimal control
376 approaches). In Fig. 3C, we show that the mean model EMG
377 predictions outperform the shuffled experimental EMG data
378 (i.e., having the same distribution as the ground truth EMG;
379 two-sided t-test, biceps $p = 2.2e-6$, triceps long head $p = 1.4e-$
380 6 , triceps lateral head $p = 4.8e-4$; P-values were Holm-
381 Bonferroni corrected for multiple comparisons, $N = 60$ shuffled
382 trials and 6 synthesized means).

383

384 **Mouse motor learning and optimal control**

385 The progression of reach kinematics and muscle patterning in
386 mice learning a novel task is a relatively understudied
387 phenomenon. We investigated the possibility that mice
388 approach an optimal motor control solution during the
389 progression of training by evaluating optimal control
390 predictions during early and late sessions of training. In Fig. 3,
391 we compared the optimal control predictions with reaches
392 selected from expert mice (i.e., after at least 5 sessions of
393 successful reaching). In Fig. 4, we compared how the optimal
394 control predictions varied when the reaches were chosen in the
395 early (i.e., in the first 3 sessions after the first successful reach
396 to pellet) or late stages of learning. We found that mice tended
397 to use muscle excitation patterns that converged more closely
398 to those derived from optimal control in the later stages of
399 learning. These results were significant when pooling the data
400 across all recorded EMG channels but not on individual
401 channels, likely because of our small sample size (early $N = 4$,
402 late $N = 6$ for means comparisons, early $N = 40$, late $N = 60$ for
403 trial-to-trial comparisons shown in Fig. 4. Comparison of trial-
404 to-trial data was compared with a two-sided t-test with a p-
405 value of $3.5e-6$).

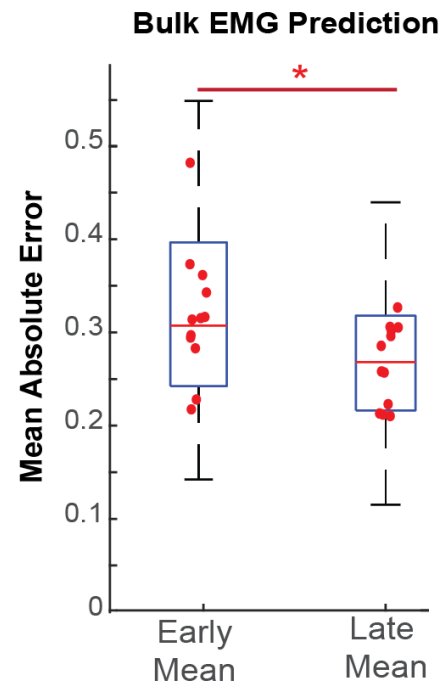


Figure 4. A comparison of the utility of the simulated muscle excitations in predicting trial-to-trial electromyography (EMG) during early or late training sessions (i.e. the first three sessions after the first successful reaches being *early* and sessions later than four sessions after the initial successful reach being *late*.) Late reaches, on bulk, have a significantly lower mean absolute error than early sessions (two-sided t-test, $p = 3.5e-6$, early $N = 40$, late $N = 60$) for predicting trial-to-trial EMG activity. Black box plots denote median (red horizontal bar), 25 to 75th percentile distributions (black box), and 10th to 90th percentile distributions (short horizontal black lines and stems).

406 **DISCUSSION**

407 Mouse models are widely used to study the neural control of movement, motor disorders, muscle
408 physiology and develop novel brain-computer interfaces and neurotechnologies. Despite the widespread
409 use of mice in the health sciences, the only available biomechanical model of the mouse forelimb is based
410 on educated guesses, which could lead to inaccurate kinematics and muscle activity predictions. We used
411 high-detail anatomical reconstruction from large scale light sheet microscopy scans to develop the first
412 physiological biomechanical model of the mouse forelimb in terms of musculoskeletal geometry and
413 muscle architecture (Mathewson et al., 2012). Dissections to determine the musculoskeletal geometry
414 would have been too challenging because of the small size of the mouse forelimb, especially in determining
415 the attachment points of the minuscule tendons of the elbow. Other imaging techniques such as microCT
416 would have likely also been adequate to produce sufficient soft tissue contrast for the reconstruction
417 (Charles et al., 2016). We then used this biomechanical model with optimal control for usage in optimal
418 control-based simulations to synthesize muscle coordination patterns that produce reaching movements that
419 match experimental kinematics. Accurately predicting muscle activity is very challenging because of the
420 infinite possible coordination patterns consistent with the tracked experimental kinematics and the high
421 physiological variance in the patterns observed in real mice (i.e., for very similar kinematics, mice often
422 use very different muscle activation strategies, some of which may be energetically costly, have high or low
423 co-contraction, be robust to disturbances, etc.). Our optimal control cost function only has terms to
424 encourage low energy (via the sum of muscle activations squared proxy) and producing kinematics
425 consistent with experimental data. Therefore, we would not expect the optimal control predictions to closely
426 match the experimental muscle activity on a reach-by-reach basis because of the high variability in the
427 experimental muscle patterning data. Nevertheless, we found that the mean optimal control muscle activity
428 predictions have strong resemblance with the mean empirical muscle activity (Fig. 3A). These results held
429 for all three recorded muscles with EMG (biceps, triceps long head and triceps lateral head). As far as we
430 know, this is the first work in any species, including humans, showing resemblance between synthesized
431 and experimental muscle activity for three-dimensional reaching movements with a biomechanical model.

432 Neuroscience experiments are sometimes limited in scope by the difficulty of simultaneous
433 recording of behavior, neurological signals, and, in some cases, muscle activity. Multisite muscle
434 recordings are often limited to a handful of accessible sites, and this limitation is exacerbated in mice, where
435 access to and implantation of many muscles is often infeasible. This model is meant to supplement
436 experiments where knowledge of muscle activity patterning could bring insight about the nature of neural
437 activity patterning. Scientists with behavioral data can extract an estimate of whole-forelimb muscle activity
438 from the model given a set of kinematics over time. Tracking of mouse kinematics has become broadly
439 accessible through the advent of pose-based tracking software like DeepLabCut, which was used in the
440 present study to monitor limb position during reaching behaviors (Mathis et al., 2018). The conjunction of
441 tracking and synthesis of full-limb muscle activations promises to expand research into behavioral control
442 significantly.

443 There are several extensions to our biomechanical model and computational tools possible for
444 future work. Our computational tools assume that no EMG is available during the experiments. If EMG is
445 collected during the experiment, the optimal control problem can be solved to predict the muscle activity
446 of muscles without EMG recordings while matching experimental EMG and kinematics data in tracking
447 simulations (Dembia et al., 2021). It is also possible to change the cost function in the optimal control
448 problem and produce predictive simulations that do not require any experimental data, including
449 kinematics. The optimal control problem could then predict the reaching kinematics and muscle activity
450 when there is a change in the task (e.g., a new pellet location) or to limb biomechanics (e.g., a weight placed
451 on the forelimb). This study is focused on the ballistic phase of reaching movements. We did not model the
452 grasping phase during the reach as we would have needed to include the muscles that control the wrist and
453 fingers in the model and simulate interaction with the pellet. One discrepancy between our simulation and

454 the empirical reach is that mice, before starting their reach, were resting on a bar, which we did not simulate
455 (and may impact the predicted muscle activity at the start of the ballistic phase).

456 Our simulations were evaluated with head-fixed mouse reaching. Using the biomechanical model
457 in free-reaching mice may be less accurate because it has more significant scapula movements, which we
458 assume to stay fixed in our model. In future work, researchers could either add a degree-of-freedom and a
459 joint motor to allow translational movement of the scapula or incorporate the muscles that control the
460 scapula as a free body in the biomechanical model (our scans available in supplementary materials should
461 help delineate these muscles). The optimal control solutions produce open-loop muscle coordination
462 patterns that are not responsive to noise or changing task or environmental constraints. It is however
463 possible to develop closed-loop controllers to track the optimized trajectory or to develop feedback
464 controllers with reinforcement learning, or introduce stochastic noise representing imprecise neural controls
465 (e.g., Van Wouwe et al., 2022).

466 We make our computational tools freely available as open-source. Users of our computational tools
467 should note that the optimal control predictions are expected to more closely resemble empirical muscle
468 activity on a mean-basis rather than on a trial-by-trial basis and carry the assumption of closely matched
469 kinematics. Furthermore, the predictions are expected to improve when mice have learned to perform the
470 task well as opposed to when mice are still in the early stages of learning; nevertheless, the model
471 predictions in the early stages of learning are still within one standard deviation of empirical results and
472 represent a significant improvement over randomized guesses from the naturalistic EMG distribution. An
473 exciting use case for our biomechanical model is to control it with artificial neural networks and relate the
474 activity in these networks with empirical neural activity from system neuroscience laboratories (Aldarondo
475 et al., 2024). Combining our computational tools and experimental data could lay the foundations for future
476 studies elucidating the principles that drive the control of movement.

477

478

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- 617

618 Table 1. Muscle origins and insertions.

Muscle Name	Origin Parent-Body	Origin Coordinates (m)	Insertion Parent-Body	Insertion Coordinates (m)
Anconeus	Humerus	[5.2e-3, 5e-4, 1.07e-2]	Ulna	[2.4e-3, -2e-4, 8.8e-3]
Anconeus, Short Head	Humerus	[3.6e-3, -9e-4, 9.1e-3]	Ulna	[3.8e-3, -6e-4, 8.3e-3]
Biceps, Long Head	Scapula	[8.5e-3, 2.3e-2, 1.2e-2]	Ulna	[2.1e-3, 4.9e-5, 8.8e-3]
Biceps, Short Head	Humerus	[9.1e-3, 1.5e-3, 1.2e-2]	Ulna	[2.1e-3, 9.8e-5, 8.8e-3]
Brachialis, Proximal Head	Humerus	[2.3e-3, 5.9e-5, 8.8e-3]	Ulna	[2.2e-3, 5.9e-5, 8.8e-3]
Brachialis, Distal Head	Humerus	[8.2e-3, 1.4e-3, 1.2e-2]	Ulna	[2.2e-3, 5.9e-5, 8.8e-3]
Brachioradialis	Humerus	[4e-3, 6.5e-4, 9.1e-3]	Radius	[-3e-3, 1.3e-3, 1.2e-2]
Deltoid, Medial	Clavicle	[9.3e-3, 5e-4, 1.3e-2]	Humerus	[5.5e-3, 1.2e-3, 1.2e-2]
Deltoid, Posterior	Scapula	[8.7e-3, 1.2e-3, 1.2e-2]	Humerus	[5.9e-3, 1.5e-3, 1.1e-2]
Flexorcarpiradialis	Humerus	[3.4e-3, -9.3e-4, 9.2e-3]	Hand	[-4e-3, 1.6e-3, 1.2e-2]
Infraspinatus	Scapula	[1.1e-2, 1.3e-3, 1.2e-2]	Humerus	[8.3e-3, 1.6e-3, 1.2e-2]
Latissimus Dorsi, Caudal	Spine*	[1.8e-2, -3e-3, 1e-2]	Humerus	[5.5e-3, 1.7e-3, 1.1e-2]
Latissimus Dorsi, Rostral	Spine*	[1.5e-2, -7.1e-4, 1.3e-2]	Humerus	[5.9e-3, 1.5e-3, 1.1e-2]
Pectoralis Major, Anterior	Rib-cage*	[9.1e-3, -1.6e-3, 1.7e-2]	Humerus	[5.3e-3, 1.4e-3, 1.2e-2]
Pectoralis Major, Posterior	Rib-cage*	[1e-2, -3.1e-3, 1.5e-2]	Humerus	[5.5e-3, 1e-3, 1.2e-2]
Pectoralis Minor, Clavicular	Clavicle	[1.1e-2, -9.6e-4, 1.3e-2]	Humerus	[5.6e-3, 1.2e-3, 1.2e-2]
Pronator Teres	Humerus	[3.6e-3, -6.7e-4, 9.3e-3]	Radius	[8.8e-4, 1.2e-3, 1e-2]
Subscapularis	Scapula	[1.3e-2, 4e-4, 1.3e-2]	Humerus	[8.2e-3, 6.2e-4, 1.2e-2]
Triceps, Long Head	Scapula	[9.8e-3, 1.2e-3, 1.2e-2]	Ulna	[4.2e-3, -5.7e-4, 8e-3]
Triceps, Lateral Head	Humerus	[8.4e-3, 1.6e-3, 1.2e-2]	Ulna	[4.1e-3, -4.5e-4, 8.3e-3]
Triceps, Medial Head	Humerus	[6.5e-3, 2.5e-4, 1.1e-2]	Ulna	[3.8e-3, -2.3e-4, 8.4e-3]

619 * Spinal and rib attachments made to Scapula fixed ground object.

620 Table 2. Muscle Parameters

Muscle Name	Max Isometric Force (N)	Optimal Fiber Length (m)	Pennation Angle (rad)	Tendon Slack Length (m)
Anconeus	0.023	0.003	0.1	0.0002
Anconeus, Short Head	0.02	0.0015	0.1	0.00015
Biceps, Long Head	0.093	0.0085	0.1	0.0002
Biceps, Short Head	0.018	0.005	0.1	0.0005
Brachialis, Proximal Head	0.066	0.007	0.1	0.0001
Brachialis, Distal Head	0.067	0.007	0.1	0.0001
Brachioradialis	0.02	0.007	0.1	0.0001
Deltoid, Medial	0.069	0.006	0.2	0.0002
Deltoid, Posterior	0.068	0.0035	0.2	0.0001
Flexorcarpiradialis	0.02	0.007	0.1	0.0001
Infraspinatus	0.065	0.003	0.2	0.0001
Latissimus Dorsi, Caudal	0.133	0.011	0.36	0.0005
Latissimus Dorsi, Rostral	0.1133	0.011	0.36	0.0005
Pectoralis Major, Anterior	0.233	0.008	0.3	0.0002
Pectoralis Major, Posterior	0.170	0.007	0.3	0.0005
Pectoralis Minor, Clavicular	0.033	0.0056	0.25	0.0002
Pronator Teres	0.02	0.003	0.1	0.0001
Subscapularis	0.34	0.005	0.2	0.0001
Triceps, Long Head	0.612	0.008	0.3	0.0007
Triceps, Lateral Head	0.125	0.007	0.17	0.0001
Triceps, Medial Head	0.16	0.004	0.2	0.0001

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