1	The Origin of Movement Biases During Reaching
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12 Abstract

13 Goal-directed movements can fail due to errors in our perceptual and motor systems. While these errors 14 may arise from random noise within these sources, they also reflect systematic motor biases that vary 15 with the location of the target. The origin of these systematic biases remains controversial. Drawing on 16 data from an extensive array of reaching tasks conducted over the past 30 years, we evaluated the merits 17 of various computational models regarding the origin of motor biases. Contrary to previous theories, we 18 show that motor biases do not arise from systematic errors associated with the sensed hand position 19 during motor planning or from the biomechanical constraints imposed during motor execution. Rather, 20 motor biases are primarily caused by a misalignment between eye-centric and the body-centric 21 representations of position. This model can account for motor biases across a wide range of contexts, 22 encompassing movements with the right versus left hand, proximal and distal effectors, visible and 23 occluded starting positions, as well as before and after sensorimotor adaptation.

24 Main

Accurate movements are crucial for everyday activities, affecting whether a glass is filled or spilled, or whether a dart hits or misses the target. Some of movement errors arise from sensorimotor noise, including visual noise regarding the location of targets and effectors^{1–3}, planning noise introduced when issuing a motor command, and neuromuscular noise when executing a movement^{4–6}.

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In addition, some of these errors arise from systematic biases that vary across the work space^{7–10}. The origin of these biases remains controversial (Fig 1a): Whereas some studies postulate that motor biases stem from systematic distortions in perception^{7,10–13}, others posit that biases originate from inaccurate motor planning and/or biomechanical constraints associated with motor execution^{14–16}. In the following section, we provide an overview of current models of systematic motor biases as well as outline a novel hypothesis, setting the stage for a re-analysis of published data and presentation of new experimental results.

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38 Starting at the input side, motor biases may arise from systematic distortions in the representation of the 39 perceived target position (Fig 1b). A prominent finding in the visual cognition literature is that the 40 remembered location of a visual stimulus is biased towards diagonal axes^{12,13,17}. That is, the reported visual 41 location of a stimulus is shifted towards the centroid of each quadrant. This perceptual bias does not 42 depend on the method of response, as this phenomenon can be observed when participants point to a 43 cued location or press a key to indicate the remembered location of a briefly presented visual target^{12,13}. 44 While this literature has emphasized that this form of perceptual bias arises from processing within visual 45 working memory, it is an open question whether it contributes to goal-directed reaching when the visual 46 target remains visible.

Another perceptual cause of motor biases stems from proprioception (Fig 1c). Systematic distortions in the perceived position of the hand^{18–20} and/or joint position^{21,22} can infleunce motor planning. For example, if the perceived starting position of the hand is leftwards of its true location, a reaching movement to a forward visual target would exhibit a rightward bias and a reaching movement to a rightward visual target would fall short⁷. A proprioceptive perceptual bias at the starting position has been reported to be the major source of bias in many previous studies.^{18–22}

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55 Whereas the preceding models have considered how distortions of visual or proprioceptive space might, 56 on their own, lead to reaching biases, motor biases could also originate from a misalignment in the 57 mapping between perceptual and motor reference frames. Based on classic theories of motor planning²³, 58 the start position and the target position are initially encoded in an eye-centric visual coordinate frame, 59 and then transformed to representations in a body-centric proprioceptive coordinate frame within which 60 the movement is planned (Fig 1d). Motor biases could arise from systematic distortions that occur during 61 this visuo-proprioceptive transformation process^{24–26}. Indeed, when participants are required to match 62 the position of their unseen hand with a visual target, systematic transformation biases are observed 63 across the workspace (Fig 1d; also see Methods)^{27,28}. In the current study, we develop a novel 64 computational model to capture how these transformation biases should result in systematic biases 65 during reaching.

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On the output side, it has been posited that reaching biases could arise from biomechanical factors that impact movement execution⁸. Specifically, movements may be biased toward trajectories that minimize inertial resistance and/or energetic costs^{15,16,29,30}. For example, minimizing energy expenditure would result in biases towards trajectories that minimize resistive forces or changes in joint angles^{14,31}. Moreover,

inaccuracies in the internal model of limb dynamics could produce systematic execution biases. For
 example, underestimating the weight of the limb would result in reaches that overshoot the target^{14,32}.

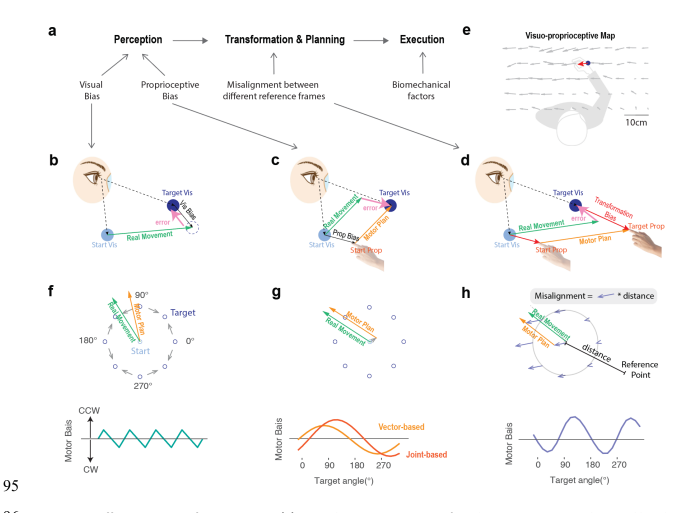
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74 To determine the origin of motor biases, we formalized four computational models to capture the 75 potential sources described above. As detailed in the Results section, the models predict distinct motor 76 bias patterns in a center-out reaching task (Fig 1f-h). While prior research has focused on the impact of 77 individual sources (e.g., vision or proprioception) on the pattern of motor errors, these studies often entail 78 a limited set of contexts (e.g., reaching behavior only when the start position is visible or only with the 79 right hand). However, looking across studies, the task context can result in dramatically different motor 80 bias patterns; indeed, when plotted in polar coordinates across the workspace, the bias functions range 81 from having single peak^{7,10,21} to quadruple peaks^{12,13,17}. This diversity underscores the absence of a 82 comprehensive explanation for motor bias phenomena across different experimental designs and setups. 83 Additionally, a notable limitation of earlier work is the reliance on small participant cohorts (n<10) and a 84 restricted number of targets (typically 8). The sensitivity of such experiments is limited in terms of their 85 capacity to discriminate between models.

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To better evaluate sources of motor bias during reaching, we report a series of experiments involving a range of contexts, designed to test predictions of the different models. We compared movements performed with the right or left hand, proximal or distal effectors, under conditions in which the start position was either visible or not visible, and before and after implicit sensorimotor adaptation. To increase the power of model comparisons, we measured the motor bias function at a higher resolution (24 targets) and in a bigger sample size (n >50 per experiment).





96 Figure 1. Different Causes of Motor Biases. (a) Motor biases may originate from biases in perceiving the initial hand 97 position (proprioceptive bias), perceiving the location of the visual target (visual bias), transforming positional 98 information from visual to proprioceptive space (transformation bias), and/or biomechanical constraints during 99 motor execution. Previous models attribute motor biases to errors originating from the distinct contributions of 100 visual (b) and proprioceptive biases (c). (d) Our model attributes motor biases to a transformation error between 101 visual and proprioceptive coordinate systems. (e) A visuo-proprioceptive map showing the matching error between 102 proprioceptive and visual space (Wang et al (2020)). Participants matched the position their hand (tip of the arrow) 103 from a random starting location to the position of a visual target (end of the arrow). The blue dot depicts an example 104 of a visual target in the workspace, and the red arrow indicates the corresponding matched hand position. 105 Participants were asked to maximize spatial accuracy rather than focus on speed. (f-h) Model bias functions

predicted by four models. Top: Illustration of how each model is applied to a center-out reaching task. As an example, the predicted motor plan and the corresponding real movement are provided for the 100° target in f and 135° target in g and h. Bottom: The predicted motor bias functions qualitatively differ in terms of the number of peaks and toughs. Note that the middle panel depicts two variants of a proprioception model.

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111 Results

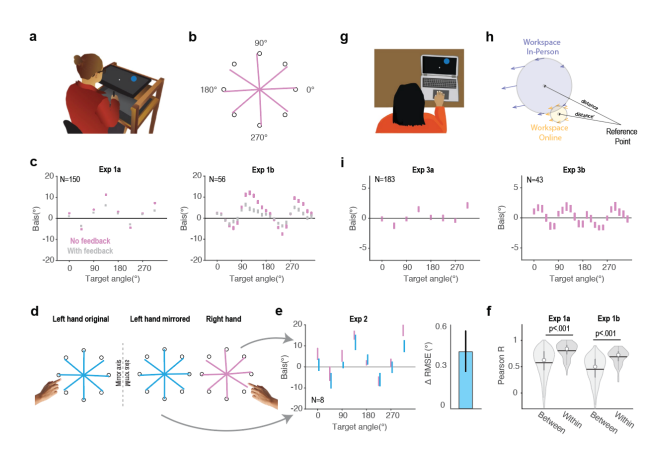
112 Motor biases across the workspace

113 To examine the pattern of motor biases during goal-directed movements, participants performed a 114 center-out reaching task with their right hand (Fig 2a). We ran two versions of the study in Experiment 1. 115 In Exp 1a, we used an 8-target version similar to that used in most previous studies^{7–10,21}. To obtain better 116 resolution of the motor bias pattern, we also conducted a 24-target version in Exp 1b. Within each 117 experiment, participants first performed the task without visual feedback to establish their baseline bias 118 and then a block with veridical continuous feedback to examine how feedback influences their biases. 119 Motor biases were calculated as the angular difference between the target and hand when the movement 120 amplitude reached the target distance (Fig 2b), with a positive error indicating a counterclockwise bias 121 and a negative error indicating a clockwise error.

122

Across the workspace, the pattern of motor biases exhibited a two-peaked function (Fig 2c) characterized by two peaks and two troughs. From the 8-target experiment, larger biases were observed for the diagonal targets (45°, 135°, 225°, 335°) compared to the cardinal targets (0°, 90°, 180°, 270°)^{33,34}. In terms of direction, reaches to diagonal targets were biased toward the vertical axis, and reaches for cardinal targets were biased in the counterclockwise direction. This pattern is similar to what has been observed in previous studies for right-handed movements^{8,9}. With the higher resolution in the 24-target experiment, we see that the peaks of the motor bias function are not strictly aligned with the diagonal targets but are

- 130 shifted towards the horizonal axis. Moreover, the upward shift of the motor bias function relative to the
- 131 horizontal line suggests that clockwise biases are more prevalent compared to counterclockwise biases
- 132 across the workspace.
- 133



135 Figure 2. Motor biases across different experimental contexts. (a) Lab-based experimental apparatus for Exps 1-2. 136 (b) Vectors linking the start position to the average endpoint position when reach amplitude equaled the target 137 radius (pink lines; Exp 1a). (c) Motor biases as a function of target location. The dots indicate the mean angular error 138 across participants during the no-feedback block (pink) and veridical feedback block (grey). The pattern of motor 139 bias was similar in Exp 1a (8-targets; left panel) and Exp 1b (24-targets; right panel), characterized by two peaks and 140 two troughs. Error bars denote standard error. (d) Motor biases generated during left hand reaches (left), left-hand 141 results when the data are mirror reversed across the vertical meridian (middle), and right-hand reaches (right). (e) 142 Left: The motor bias generated by right-hand reaches was similar to that of mirror-reversed left-hand reaches. Right: 143 Difference in RMSE when the right-hand map is compared to the original left-hand map relative to when the right-

144 hand map is compared to the mirror reversed left-hand map. Positive values indicate better data alignment when 145 the left hand data are mirror-reversed. (f) Correlation of the motor bias function between the no-feedback and 146 feedback blocks is higher in the within-participant condition compared to the between-participant condition. Gray 147 bars indicate the 25% and 75% quartiles. White dots indicate the median and horizontal lines indicate the mean. (g) 148 Experimental setup for Exp 3. Participants were asked to make center-out reaching movements using a trackpad or 149 mouse. These movements predominantly involve finger and wrist movements. (h) The workspace is presumed to be 150 closer to the reference point (e.g., left shoulder) for finger/wrist movements (Exp 3) compared to that of arm 151 movements (Exp 1-2). (i) The pattern of motor biases in finger/wrist movements for 8-targets (left) and 24-targets 152 (right).

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154 Motor biases primarily emerge from a misalignment between visual and hand reference frames

We developed a series of models to capture how systematic distortions at different sensorimotor processing stages would cause systematic motor biases (Fig 3a). Here we consider processing associated with the perceived position of the target, the perceived position of the arm/hand, and planning processes required to transform a target defined in visual space to a movement defined in arm/proprioceptive space. Biases could also arise from biomechanical constraints. Given that biomechanical biases are not easily simulated, we will evaluate this hypothesis experimentally (see below).

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We implemented four single-source models to simulate the pattern of the motor bias that would be predicted in a center-out reaching task (Fig 1f-h; see Methods). For the Visual Bias model, we assumed that the representation of the visual targets were biased towards the diagonals¹³. This model predicts a four-peaked motor bias function (Fig 1f). For the Proprioceptive Bias model, we considered two variants building on the core idea is that the perceived starting position of the hand is distorted: A Vector-Based model in which the motor plan is a vector pointing from the perceived hand position to the target^{7,10} and a Joint-Based model in which the movement is encoded as changes in the shoulder and elbow joint angles to move the limb from a start position to a desired target location^{21,22} (See Fig S1). Importantly, both models predict a motor bias function with a single peak (Fig 1g). Taken together, models that focus on systematic distortions of perceptual information do not qualitatively capture the observed two-peaked motor bias function (Fig 2c).

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174 The fourth model, the Transformation model is based on the idea that the start and target positions are 175 initially encoded in visual space and transformed into proprioceptive space for motor planning²³. Motor 176 biases may arise from a transformation error between coordinate systems. Two prominent features stand out when this transformation error is empirically measured^{18-20,27}. First, the direction of the 177 178 transformation error is similar across the workspace (e.g., a leftward and downward matching error for 179 right-handers). Second, the magnitude of the error increases with distance from the body (Fig 1d). As such, 180 we simulated a visuo-proprioceptive error map by using a leftward and downward error vector with the 181 magnitude scaled across the workspace based on the distance of each location to a reference point (Fig 182 1h Top). This model predicts a two-peaked motor bias function (Fig 1h Bottom), a shape strikingly similar 183 to that observed in Exps 1a and 1b.

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185 To quantitively compare the models, we fit each model with the data in Exp 1b. The Transformation Bias 186 model provided a good fit of the two-peaked motor bias function (R²=0.84, Fig 3a left, see Table S1 for 187 parameters). Fig 1h (top) shows the recovered visual-proprioceptive bias map based on the parameters 188 of the Transformation Bias model when fit to the reaching data in Exp 1b. The simulated results are very 189 similar to the map measured empirically in the previous study²⁰ (Fig 1e). In contrast, the Visual Bias and 190 Proprioceptive Bias models provide poor fits (all R²<0.18, Fig 3a right). Thus, the simulations suggest that 191 motor biases observed in reaching primarily originate from a transformation between visual and 192 proprioceptive space.

193

194 A second way to evaluate the models is to compare the motor bias functions for the left and right hands. 195 The Proprioceptive and Transformation Bias models predict that the bias function will be mirror-reversed 196 for the two hands whereas the Visual Bias model predicts that the functions will be superimposed. We 197 compared the functions for right and left hand reaches in Exp 2 using the 8-target layout. We found that 198 the dissimilarity (RMSE) between the pattern of motor biases across two hands significantly decreased 199 when the left-hand data are mirror-reversed compared to when the bias patterns are compared without 200 mirror reversal (t(78) = 2.7, p = 0.008, Fig 2d-e). These results are consistent with the Transformation Bias 201 model and provide further confirmation that the Visual Bias model, at least as conceptualized here, fails 202 to provide a comprehensive account of reaching biases. 203 204 While the overall pattern of biases in the visuo-proprioceptive map are similar across individuals, there 205 are subtle individual differences^{18,20}. As such, we would anticipate that the motor bias function would also 206 exhibit stable individual differences due to idiosyncrasies in the visuo-proprioceptive map. To examine 207 this, we correlated the bias functions obtained from blocks in which we either provided no visual feedback 208 or veridical endpoint feedback. The magnitude of the biases was attenuated when endpoint feedback was

provided, likely because the feedback reduced the visuo-proprioceptive mismatch. Nonetheless, the overall pattern of motor bias was largely preserved, with the within-participant correlations (Exp 1a: $r_{norm} = 0.999$, Exp 1b: $r_{norm} = 0.974$) significantly higher than the averaged between-participant

212 correlation in both Exp 1a and Exp 1b (Fig 2f).

213

214 <u>Visual bias also contributes to the motor bias.</u>

In the preceding section, we considered each model in isolation, testing the idea that motor biases arisefrom a single source. However, the bias might originate from multiple sources. For example, there could

be a distortion in both vision and proprioception, or a visuo-proprioceptive transformation that operates on distorted inputs. To address this, we evaluated hybrid models by combining the Visual Bias model with the Proprioceptive or Transformation Bias models. Although theoretically plausible, we did not consider a hybrid of the Proprioceptive and Transformation Bias models since they conflict in terms of whether the start position is perceived visually or proprioceptively.

222

223 The hybrid model that combines the Transformation and Visual Bias models (T+V model) provided an 224 excellent fit of the motor bias pattern in Exp 1b (R²=0.973, Fig 3a). Based on a comparison of BIC values, 225 this model not only outperforms the other hybrid models, but also significantly improved the fit compared 226 to the Transformation Bias model alone. These results are especially interesting in that the assumed visual 227 bias towards the diagonal axes has only been shown in studies in which perception was tested after the 228 target had been extinguished. The current results suggest that this bias is also operative when the target 229 remains visible, suggesting that the visual bias may reflect a general distortion in how space is represented, 230 rather than a distortion that arises as information is processed in visual working memory.

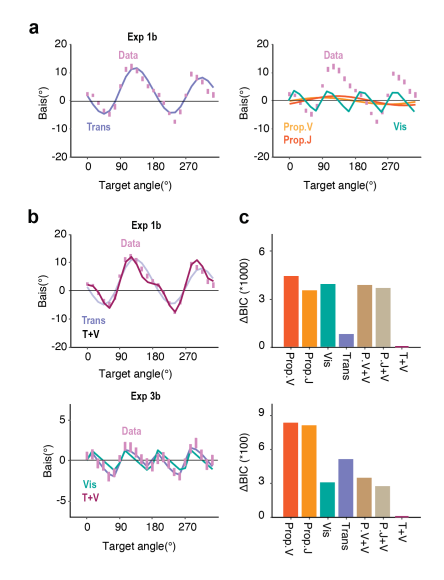


Figure 3. The pattern of motor biases is best explained by assuming systematic distortions in the perceived location of the target and the transformation between visual-proprioceptive coordinate frames. (a) For singlesource models, the pattern of motor biases in the no feedback block of Exp 1a (pink dots) is best fit by the Transformation Bias model (left) compared to the other models (right). (b) A mixed model with transformation and visual biases (T+V) provides the best fit to the motor bias function in both Exp 1b (top) and Exp 3b (bottom). (c) Model comparison using BIC. ΔBIC values are provided by subtracting the BIC from the best performing model (i.e., the T+V model). A smaller ΔBIC signifies better model performance.

239

To further evaluate the T+V model, we examined its performance in explaining the motor bias function obtained in an on-line study (Exp 3) in which participants performed the center-out task by moving a finger across a trackpad. One major difference between the in-person and on-line setups is that the workspace is much smaller and closer to the body when participants use a trackpad (Fig 2g). As such, the magnitude of the motor biases generated by transformation errors should be smaller in the online compared to the in-person setup (Fig 2h).

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247 Consistent with the prediction of the T+V model, we found markedly smaller motor biases with the online 248 setup (Exp 3) compared to the in-person setup (Exp 1) (Fig 2c). While the motor bias functions were similar 249 across experiments, we observed two small peaks between 20° and 200° in Exp 3 that were not apparent 250 in Exp 1. When we fit this function to single source models, the Visual Bias model outperformed the 251 Transformation Bias model. This suggests that, when the movements are close to the body, visual biases 252 make a relatively stronger contribution to the motor biases compared to transformation biases. 253 Nonetheless, the T+V model again provides the best fit to the motor bias function (R²=0.857, Fig 3e, see 254 Table S1 for parameters), significantly outperforming the other alternatives including the Visual Bias 255 model (Fig 3f).

256

257 <u>Transformation model accounts for qualitative changes in the motor bias function</u>

The Transformation Bias model assumes that, for normal reaching, both the start and the target positions are encoded in visual space before being transformed into proprioceptive space for motor planning. However, if the start position is not visible, then the sensed start position would be directly encoded in proprioceptive space (i.e., where the hand is positioned), bypassing the need for a transformation between coordinate frames. As such, biases arising from the transformation process would only arise when the input is limited to the perceived position of the visual target. When we simulated the scenario

- in which the start position is not visible, the Transformation Bias model predicts a single-peaked function
- 265 (Fig 4a right), a qualitative change from the two-peaked function predicted when both the start position
- and target position are visible (Fig 4a left).
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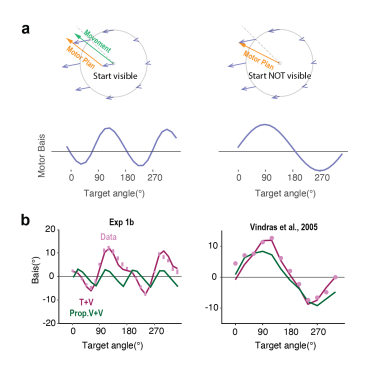


Figure 4. Motor bias patten changes when the start position is not visible. (a) Schematic showing the planned movement under the Transformation Bias model when the start position is either visible (left) or not visible (right). In the latter case, only the target position has to be transformed from visual to proprioceptive coordinates with the start position directly encoded in proprioceptive space. The T+V model now predicts a single-peaked motor bias function (lower row). (b) Consistent with this prediction, a two-peaked function is predicted when the start position is visible (as in Exp 1) and a single-peaked function is predicted when start position is not displayed. Data (pink dots) are from Vindras et al (2005).

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268

To test this idea, we re-examined data from previous studies in which the participant's hand was passively moved to a start position with no visual information given about the start location or hand position^{7,10,21}. Strikingly, the motor bias function under this condition has only have one peak. Thus, the transformation

Bias model provides a novel account of the difference in motor biases observed when the start position
is visible (Exp 1-3) compared to when it is not visible.

282

283 We note that the one-peaked motor bias function has previously been interpreted as evidence in support 284 of a Proprioceptive Bias model (Fig 1f)^{7,10,21}. We performed a model comparison on the data from one of 285 these studies¹⁰ and the T+V outperformed the Proprioceptive Bias, as well as the P+V models (Δ BIC=10.9). 286 In addition, only the T+P model is able to account for the asymmetry between clockwise and 287 counterclockwise biases. In summary, these results suggest that motor biases when reaching from an 288 unseen start position arise when the target position is transformed from visual to proprioceptive 289 coordinates rather than from a proprioceptive bias impacting the sensed start position. Moreover, the 290 T+V model provides a parsimonious account of the bias functions, independent of the visibility of the start 291 position.

292

Another way to evaluate the Transformation model is to perturb the position of the visual start position relative to the real hand position. Under this manipulation, a single peaked motor bias function is observed (Fig S2)^{21,22}. Interestingly, the functions exhibit opposing phase shifts when the starting position is perturbed to the left versus to the right (Fig S2). This qualitative change in the motor bias function can again be successfully captured by the Transformation model (Fig S2c). Taken together, these data provide strong evidence favoring the notion that motor biases originate primarily from a misalignment between visuo-proprioceptive reference frames.

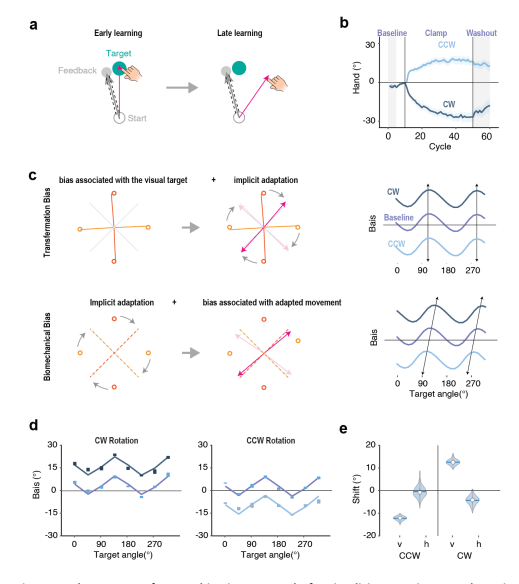
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301 <u>Biomechanical models fail to account for motor biases</u>

302 An alternative account of motor biases is that they arise from biomechanical constraints associated with 303 upper limb movements. For example, movement kinematics have been explained in terms of cost functions to minimize energy and/or minimize jerk,³⁵ constraints that may result in an increase in endpoint error^{15,16,29}. One argument against a biomechanical model comes from evidence provided in the previous section: Biomechanical constraints associated with movement execution would not predict qualitative changes in the motor bias function in response to a visual manipulation of the start position.

308

309 As a second comparison of the T+V and Biomechanical Bias models, we examined how motor biases 310 change after the sensorimotor map is recalibrated following a form of motor learning, implicit 311 sensorimotor adaptation. Here we re-analyzed the data from previous experiments that had used a 312 perturbation technique in which the visual feedback was always rotated by 15° from the target, 313 independent of the hand position (Fig 5a, non-contingent clamped feedback³⁶). Participants adapt to this 314 perturbation, with subsequent reaches to the same target shifted in the opposite direction (Fig 5b), 315 reaching an asymptote of around 20° and showing a robust aftereffect when the perturbation is removed. 316 Participants are unaware of their change in hand angle in response to clamped feedback, reporting their 317 perceived hand position to be close to the target³⁷.



319 Figure 5. The pattern of motor bias is preserved after implicit sensorimotor adaptation, consistent with the 320 Transformation + Visual Bias model. (a) Illustration of the clamped perturbation. Feedback cursor is offset by a fixed 321 angle from the target, independent of the participant's heading direction. (b) Time course of hand angle in response 322 to clockwise or counterclockwise clamped feedback. Vertical lines demarcate the perturbation block which was 323 preceded and followed by no-feedback baseline and washout phases, respectively (gray areas). Shaded area 324 indicates standard error. (c) Predictions for the bias functions after adaptation for the T+V (top) and Biomechanical 325 models (bottom). See text for details. The right column shows the predicted motor bias functions following 326 adaptation in response to a clockwise (CW) or counterclockwise (CCW) clamp. (d) Motor bias functions before and 327 after training in a CW (left) and a CCW (right) clamp. Data taken from Morehead et al. (2017) and Kim et al. (2018);

328	the height of the colored bars indicates the standard error for each data point. The best-fit lines for the T+V model
329	are shown. (e) Parameter values to capture vertical and horizontal shifts in motor bias functions before and after
330	training. The CW and CCW conditions both showed a significant vertical shift but no horizontal shift.
331	

For the T+V model, the transformation between visual and proprioceptive space depends on the perceived positions of the start and target locations in a visual-based reference space, one that remains unchanged before and after adaptation. We assume that adaptation has changed a sensorimotor map that is referenced after the transformation from visual to proprioceptive space. As such, the heading angle after adaptation for each target location is obtained by summing the motor biases for that target location and the extent of implicit adaptation. This would result in a vertical shift of the motor bias function (Fig 5c top).

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In contrast, the biomechanical model predicts that motor biases will be dependent on the actual movement direction rather than the target location (e.g., a bias towards a movement that is energetically efficient). Since the mapping between a target location and its corresponding reach direction is rotated after adaptation, the motor bias pattern would also be rotated (Fig 5c bottom). As such, the biomechanical model predicts that the motor bias function will be shifted along both the horizontal and vertical axes.

345

To arbitrate between these models, we analyzed the data from two previous studies, looking at the bias function from no-feedback trials performed before (baseline) and after adaptation (washout) ^{36,38}. Consistent with the prediction of the T+V model, the motor bias function shifted vertically after adaptation (Fig 5d) but did not shift horizontally.

351 To quantitatively evaluate these results, we first fit the motor bias function during the baseline phase with 352 the T+V model and fixed the parameters. We then examined the heading angles during the aftereffect 353 phase by fitting two additional parameters, one that allowed the function to shift vertically (v) and the 354 other to allow the function to shift horizontally (h). The T+V model predicts that only v will be different 355 than zero; in contrast, the Biomechanical model predicts that h and v will both be different than zero and 356 should be of similar magnitude. The results clearly favored the T+V model (Fig 5e). The vertical shift in the 357 bias functions was of a similar magnitude as the aftereffect, with the shift direction depending on the 358 direction of the clamped feedback (v: CW: 12.5°; CCW: -12.2°, p<0.001). In contrast, the best fitting value 359 for h was not significantly different from zero in both condition (Fig 5g). These results are consistent with 360 the hypothesis that visual representations are first transformed into proprioceptive space for motor 361 planning, with the recalibrated sensorimotor map altering the trajectory selected to achieve the desired 362 movement outcome.

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- 364

365 Discussion

366 While motor biases are ubiquitous in goal-directed reaching movements, the origin of these biases has 367 been a subject of considerable debate. We addressed this issue by characterizing these biases across a 368 range of experimental conditions, and evaluated a set of computational models derived to capture 369 different possible sources of bias. Contrary to previous theories, our results indicate that motor biases do 370 not stem from a distortion in the sensed position of the hand^{7,10,21,22} or from biomechanical constraints 371 during movement execution^{15,16,29}. Instead, motor biases appear to arise from systematic distortions in 372 perceiving the location of the visual target and the transformation required to translate a perceived visual target into a movement described in proprioceptive coordinates.^{24–26} Strikingly, our model successfully 373 374 accounts for sensorimotor biases across a wide range of contexts, encompassing movements performed

with either hand as well as with proximal and distal effectors. Our model also accounts for the qualitative changes in the motor bias function that are observed when vision of the starting position of the hand is occluded, and when the sensorimotor map is perturbed following implicit adaptation.

378

379 While motor biases have been hypothesized to reflect a mismatch across perceptual and motor 380 coordinate systems,^{25,26} it is unclear what information is transformed and what reference frame is 381 employed for motor planning. Interestingly, many previous studies posit that movement is planned in an 382 eye-centric visual reference frame³⁹⁻⁴¹. While the target can be directly perceived in this reference space, 383 the start position of the hand would need to be transformed from a proprioceptive reference frame to a 384 visual one. Systematic error in this transformation would mean that the start position of the hand is 385 inaccurately represented in visual space, resulting in motor biases^{7,25}. This idea underlies the 386 Proprioceptive Bias models described in this paper.

387

In contrast to these models, our Transformation Bias model posits that movement is planned in a handcentric proprioceptive reference frame. By this view, when both the target and start position are provided in visual coordinates, the sensorimotor system transforms these positions from visual space to proprioceptive space. Systematic error in this transformation process will result in motor biases. When vision of the start position is available, the Transformation Bias model successfully accounts for the twopeaked motor bias function (Exp 1), whereas the Proprioceptive Bias model in which motor planning is eye-centric predicts only a single peak in the function.

395

396 Perhaps most compelling, the Transformation Bias model accounts for how the pattern of motor biases 397 change when the visibility of the start position is manipulated. When visible, the motor bias function is 398 two-peaked. In contrast, when the start position is occluded, the transformation from visual to

399 proprioceptive space is only relevant for the target position since the start position of the hand is already 400 represented in proprioceptive space. Here the model predicts a motor bias function with a single peak, a 401 function that has been observed in previous studies^{7,10}.

402

We note that there is a third scenario, one which both the start position and target position are provided in proprioceptive space. We predict that under this condition, motor biases originating from the visuoproprioceptive transformation would completely disappear. Indeed, when the hand is passively moved first to the target location and then to the start position, subsequent reaches to the target do not show the signature of bias from a visuo-proprioceptive transformation.¹² Instead, the reaches exhibited a bias towards the diagonal axes, consistent with the predicted pattern if the sole source of bias is visual.

409

410 Why would a sensorimotor system exhibit inherent biases during the transformation process? We 411 propose that these biases arise from two interrelated factors. First, these systems are optimally tuned for 412 distinct purposes: A body-centric system predominantly uses proprioceptive and vestibular inputs to 413 determine the orientation and position of the body in space, while an eye-centric system relies on visual 414 inputs to interpret the layout of objects in the external world, representations that should remain stable 415 even as the agent moves about in this environment.^{42,43} Second, these sensory systems consistently 416 receive information with very different statistical distributions^{2,44}, perhaps because of these distinct 417 functions. For example, visual inputs tend to cluster around the principal axes (horizontal and vertical) 418 ^{45,46}, whereas proprioceptive information during reaching is clustered around diagonal axes⁴⁷. This is 419 because these movements are often the least effortful and are the most frequently enacted directions of 420 movement.⁴⁸. The differences in computational goals and input distributions might have led to natural 421 divergences in how each system represents space⁴⁹, and consequently, result in a misalignment between 422 the reference frames.

Λ	2	2
+	7	2

The Transformation Bias model addresses how biases arise when the information is passed along from a visual to a proprioceptive reference frame. However, the results indicate that another source of bias originates from a distortion within the visual reference frame itself, manifesting as an attractive bias towards the diagonal axes. Thus, the best fitting model posits two sources of bias, one related to the representation of the visual target and a second associated with the transformation process. This hybrid Transformation + Visual Bias model outperformed all single-source and hybrid models, providing an excellent fit of the behavioral data across a wide variety of contexts.

431

432 There are several hypotheses concerning the origin of this bias. One account has focused on the idea that 433 these biases arise from distortions introduced in visual working memory. The biases are observed when 434 participants report the location of a remembered visual target^{13,17,50}, independent of the reporting 435 method (pointing or keypresses)¹⁷. However, as mentioned above, similar biases are observed even when 436 participants report the location of a proprioceptive target (i.e., participants match the position of their 437 unseen hand).¹² Moreover, as shown in our study, diagonal target biases make a sizable contribution even 438 when the visual target is always present, imposing no demands on working memory. As such, we postulate 439 that the visual biases may reflect a more domain-general distortion of spatial representations.

440

Our data suggest that biomechanical factors do not significantly impact motor biases. While we did not formalize a biomechanical model, we provided several lines of evidence suggesting the biomechanical factors have minimal influence on the pattern of motor biases. For example, it is hard to envision a biomechanical model that would account for the qualitative change in the bias function when the start position was visible (two-peak function) to when it was hidden (one-peak function).

447 Empirically, we evaluated biomechanical contributions to motor biases by examining the bias pattern 448 observed before and after implicit sensorimotor adaptation. We assume that adaptation mainly modifies 449 a sensorimotor map⁵¹ but has a relatively smaller influence on a visuo-proprioceptive map^{28,37,52}. That is, 450 adaptation may change the mapping between a target represented in the proprioceptive space and the 451 motor commands required to reach that location. Given that a biomechanical model assumes that motor 452 biases are associated with the direction of a movement, this model would predict that the pattern of 453 motor biases would be distorted by implicit motor adaptation. At odds with this prediction, the pattern 454 of motor biases remained unchanged after adaptation, a result consistent with the Transformation Bias 455 model.

456

Nonetheless, the current study does not rule out the possibility that biomechanical factors may cause motor biases in different contexts. In our study, biomechanical constraints may not be influential since the extent of the required movements was relatively modest and involved minimal interaction torques. Moreover, we focused on examining biases that manifest at the movement endpoint rather than in the movement trajectory where biomechanics are known to play a greater role.^{15,16} Future studies are necessary to explore whether biomechanical biases are more pronounced in contexts in which the limb dynamics are crucial and/or the movements are energetically costly.

464

465 Methods

466 <u>Participants</u>

For the lab-based study (Exp 1, 2), 206 undergraduate students (age: 18-24) were recruited from University of California, Berkeley. For the online study (Exp 3), 183 young adult participants (age: 18-30) were recruited via Prolific, a website designed to recruit participants for online behavioral testing. All participants were right-handed as assessed by the Edinburgh handedness test⁵³ with normal or corrected471 to-normal vision. Each participant was paid \$15/h. The protocol was approved by the institutional review

- 472 board at the University of California Berkeley.
- 473

474 <u>Procedure</u>

475 Experiments 1a, 1b, and 2:

Experiments 1a, 1b, and 2 were conducted in the lab. Participants performed a center-out reaching task,
holding a digitizing pen in the right or left hand to make horizontal movements on a digitizing tablet
(49.3cm x 32.7cm, sampling rate= 100 Hz; Wacom, Vancouver, WA). The stimuli were displayed on a 120
Hz, 17-in. monitor (Planar Systems, Hillsboro, OR), which was mounted horizontally above the tablet (25
cm), to preclude vision of the limb. The experiment was controlled by custom software coded in MATLAB
(The MathWorks, Natick, MA), using Psychtoolbox extensions, and run on a Dell OptiPlex 7040 computer
(Dell, Round Rock, TX) with Windows 7 operating system (Microsoft Co., Redmond, WA).

483

Participants made reaches from the center of the workspace to targets positioned at a radial distance of 8 cm. The start position and target location were indicated by a white annulus (1.2 cm diameter) and a filled blue circle (1.6 cm), respectively. Vision of the hand was occluded by the monitor, and the lights were extinguished in the room to minimize peripheral vision of the arm. Feedback, when provided, was in the form of a 4 mm white cursor that appeared on the computer monitor, aligned with the position of the digitizing pen.

490

To start each trial, the participant moved the cursor to the start circle (5mm diameter). After maintaining the cursor within the start circle for 500 ms, a target appeared at one of the target locations. The participant was instructed to make a rapid slicing movement through the target. We did not impose any reaction time guidelines, allowing the participant to set their own pace to initiate the movement. On no-

495	feedback trials, the cursor was blanked when the hand left the start circle, and the target was extinguished
496	once the radial distance of the movement reached the target distance (8 cm). On feedback trials, the
497	cursor was visible throughout the movement until the movement amplitude reached 8 cm; at that point,
498	its position was frozen for 1 s, providing feedback of the accuracy of the movement (angular position with
499	respect to the target). After this interval, the target and cursor were extinguished.
500	
501	At the end of both the no-feedback and feedback trials, a white ring appeared denoting the participant's
502	radial distance from the start position. This ring was displayed to guide the participant back to the start

position without providing angular information about hand position. Once the participant moved within 2 cm of the start position, the ring was extinguished, and a veridical cursor appeared to allow the participant to move their hand to the start position. Participants had to remain within the start position for 200 ms before the next target appeared. If the amplitude of the hand movement did not reach the target (<8 cm radial distance) within 300 ms, the message "too slow" would be displayed for 500 ms before the white ring appeared.

509

510 For Exp 1a and Exp 2, there were 8 target locations, evenly spaced in 45° increments around the workspace 511 (primary axes and main diagonals). For Exp 1b, there were 24 target locations, evenly spaced in 15° 512 increments. Each experiment consisted of a no-feedback block followed by a feedback block. There were 513 5 trials per target (40 trials total) for each block in the Exps 1a. There were 4 trials per target (96 trials 514 total) in Exp 1b.

515

516 Experiments 3a and 3b

517 Exps 3a and 3b were conducted using our web-based experimental platform (Tsay et al., 2021). 518 Participants made center-out movements by controlling a cursor with the trackpad on their personal

computers. It was not possible to occlude vision of the hand. However, since the visual stimulus was presented on a vertical monitor and the hand movement was in the horizontal plane, we assume vision of the hand was limited to the periphery (based on observations that the eyes remain directed to the screen during the trial). The size and position of visual stimuli were scaled based on each participant's screen size (height = 239.6 \pm 37.7 mm, width = 403.9 \pm 69.5 mm). The experiment was controlled by custom software written with JavaScript and presented on Google Chrome. Data were collected and stored using Google Firebase.

526

527 The procedure was designed to mimic the lab-based experiments. On each trial, the participant made a 528 center-out planar movement from the start position to a visual target. A white annulus (1% of screen 529 height in diameter, 0.4 cm on average) indicated the start position, and a blue circle (1% of screen height 530 in diameter) indicated the target location. The radial distance of the target from the start position was 40% 531 of the screen height (5 cm on average). At the beginning of each trial, participants moved the cursor (0.6% 532 of the screen height in diameter) to the start position, located at the center of their screen. The cursor 533 was only visible when its distance from the start position was within 20% of the screen height. After 534 maintaining the cursor at the start position for 500 ms, the target appeared. The participant made a rapid 535 slicing movement through the blue target. As in the online experiments, there were feedback and no-536 feedback trials. For feedback trials, the cursor was visible until it reached the target distance, and then 537 froze for 1 s at the target distance. On no-feedback trials, the cursor was extinguished after the hand 538 exited the start position and the target disappeared once the radial distance of the movement reached 539 the target distance. 500 ms after the end of the trial, the cursor became visible, repositioned at a random 540 location within 10% of the screen height from the start position. The participant then moved the cursor 541 to the start position to trigger the next trial.

There were 8 target locations in Exp 3a and 24 target locations in Exp 3b. As with the lab-based experiments, each experiment included a no-feedback block followed by a feedback block. We obtained larger data sets in the online studies: For each block, there were 20 trials/target (160 total trials for Exp 3a and 480 total trials for Exp 3b).

547

548 *Reanalysis of prior data sets*

Vindras et al (2005). This study used a design in which the participant did not see the start position of the movement. This was achieved by not included start position information in the visual display and passively moving the participant's hand to a start position prior to each reach. Once positioned, a visual target would appear and the participant reached to that location. Across trials, there were two start positions, 12 target positions (spaced evenly by 30° around the workspace), and two target distances (6 and 12 cm). In modeling these data, we used the movement endpoint averaged across start positions and target distances.

556

557 Morehead et al (2017) & Kim et al (2018). We re-analyzed the data from the 15° conditions of Exp 4 in 558 Morehead et al (2017) and Exps 1 and 2 in Kim et al (2018). These three experiments examined visuomotor 559 adaptation using non-contingent clamped feedback. On perturbation trials, the feedback cursor was 560 presented at the radial position of the hand but with a fixed 15° angular offset relative to the target. 561 Participants were informed that the angular position was not contingent on their hand position and 562 instructed to move directly to the target, ignoring the feedback. This method results in robust implicit 563 adaptation, with the heading direction of the movement gradually shifting away from the target in the 564 opposite direction of the cursor. Participants are unaware of this change in behavior (Tsay et al, 2020). In 565 each experiment, there were three blocks: A no-feedback baseline block (10 trials/target), a clamped 566 feedback block (60 trials/target), and a no-feedback washout block (10 trials/target).

567

568 Data analyses

569 Motor bias refers to the angular difference between the position of the hand and target when the hand

570 reaches the endpoint target distance. Angular errors were plotted as a function of the target position with

- 571 0° corresponding to the rightward target (3 o'clock location) and 90° corresponding to the forward target.
- 572 Positive bias values indicate a counterclockwise error, and negative values indicate a clockwise error.

573

To assess the similarity of the motor bias functions across different conditions, we calculated the normalized correlation coefficient as $r_{norm} = \frac{r_{data}}{r_{max}} \cdot r_{data}$ is the Pearson correlation coefficient between the two motor bias functions. r_{max} is the correlation coefficient between the recorded motor bias function and the true (but unknown) underlying motor bias function from that condition. To calculate r_{max} , we used a method developed to measure the noise ceiling for EEG/fMRI data ⁵⁴:

579
$$r_{max} = \frac{2}{1 + \sqrt{\frac{1}{r_{half}^2}}}$$

where r_{half} is determined by splitting the data set (based on participants) into random halves and calculating the correlation coefficient between the first half and the second half of the data. We bootstrapped r_{half} by resampling the data 2000 times and used the average value. r_{max} is calculated separately for a pair of conditions and the smaller one is applied as the normalizer for r_{norm} .

584

585 <u>Models</u>

To examine the source of motor bias, we considered five single-source models and three multiple-sourcemodels.

589 Visual Bias model

590 The Visual Bias model postulates that movement biases arise because the perceived position of the visual 591 target is systematically distorted (Fig 1b). Here we draw on the work of Huttenlocher et al¹³. In their study, 592 a visual target was picked from an invisible circle, presented for 1 s and then blanked. The participant then 593 indicated the remembered position of the target by pointing to a position on a circular digitizing pad. The 594 results showed a bias towards the four diagonal directions (45°, 135°, 225°, 315°), with the magnitude of 595 this bias increasing linearly as a function of the distance from the diagonals. As such, the maximum bias 596 was observed for targets close to four cardinal target locations (0°, 90°, 180°, 270°), and the sign of bias 597 flipped at the four cardinal target locations.

598

We used the shape of this function to model bias associated with the perception of the location of the visual targets. To obtain a continuous function, we assumed a transition zone around the cardinal targets, each with a half-width represented by the parameter *a* (Fig 1a), and the peak motor bias is represented

by the parameter *b*. As such, the angular bias (y) at a target located at x° can be formalized as:

603 $[1] x' = x \mod(90)$

604 [2]
$$y = \frac{b * x'}{a}$$
, if $x' < a$;

605
$$y = b - \frac{(x'-a)b}{45^\circ - a}$$
, if $90 - a > x' > a$;

606
$$y = -\frac{b * (90 - x')}{a}, if x' > 90 - a;$$

607 This model has two free parameters (*a* and *b*). If participants directly reach to the perceived target location,

608 their motor biases will directly reflect their visual biases.

609

610 Vector-based Proprioceptive Bias model

611 Vindras et al. ^{7,10} proposed a model in which movement biases result from a misperception in estimating 612 the initial position of the hand (Fig 1c). Specifically, it has been shown that the perceived position of the 613 hand when placed near the center of the workspace is biased towards the ipsilateral side and away from the body ^{18,19,27}). Assuming that the planned movement is formed by a vector pointing from the sensed 614 615 hand position to the visual target position, this proprioceptive distortion will result in systematic motor 616 biases around the workspace. For example, for the target at 90°, misperceiving the initial position of the 617 right hand to the right of the start position will result in a movement that is biased in the counterclockwise 618 (leftward) direction.

619

To simulate this Proprioceptive Bias model, we assumed the participants perceived the start position (0, 0) as a rightward bias away from the midline position, defining a proprioceptive error vector (x_e, y_e) . For a target i at $[x_i, y_i]$, the motor plan is a vector $[x_i - x_e, y_i - y_e]$. From this, we calculated the angular difference between the motor plan vector and the target position to generate the motor bias for each target. The two free parameters in this model are $[x_e, y_e]$.

625

626 Joint-based Proprioceptive Bias model

Reaching movements may also be planned in joint coordinates rather than the hand (endpoint) position 21,22 . Based on this hypothesis, motor biases could come about if there is a misperception of the initial elbow and shoulder joint angles. To implement a Joint-Based Proprioceptive Bias model, we represent the length of the forearm and upper arm as l1 and l1, respectively. We denote the initial angles of the shoulder and elbow joints as θ_0 and φ_0 , respectively, and their associated perceived error as θ_e , and φ_e (See Fig S1).

634 By setting the origin of the coordinate system for the right shoulder at P_0 (0, 0), the hand can be 635 represented as: 636 $[3] P(\theta, \varphi) = [l1\cos\theta_0 + l1\cos\varphi_0, l1\sin\theta_0 + l2\sin\varphi_0].$ 637 For a fixed position in the workspace, there will be a unique solution pair for θ and $\varphi(\pi > \varphi > \theta > 0)$, 638 should a solution exist. To calculate the required change in joint angle to reach a visual target, we assumed 639 that the system plans a movement based on the perceived hand position: 640 [4] $P_p = P(\theta_0 + \theta_e, \varphi_0 + \varphi_e)$ Then we solve the following equation to decide the proper $\sigma \theta_i$ and $\sigma \phi_i$ that transfer the hand from the 641 642 start position to a target i at $[x_i, y_i]$: 643 $[5] P_p - [x_i, y_i] = P(\theta_0 + \theta_e + \sigma \theta_i, \varphi_0 + \varphi_e + \sigma \varphi_i) - P_p$ 644 We calculated the real movement direction based on the real hand position: 645 [6] $\sigma h_i = P_p - P(\theta_0, \varphi_0)$ 646 We compare the direction of σh_i and the target direction to calculate the motor bias. For simplicity, we 647 assume l1=l2=24 cm.⁵⁵ The four free parameters in this model are θ_0 , φ_0 , θ_e , and φ_e 648 649 Transformation Bias model 650 The Transformation Bias model proposes attributes motor biases to systematic errors that arise during 651 the transformation from a visual to proprioceptive-based reference frame. To implement this model, we 652 refer to an empirically derived visuo-proprioceptive error map from a data set that sampled most of 653 reachable space (Fig 1d, ²⁰). Specifically, in that study, participants were asked to move their unseen hand 654 from a random start position to a visual target. Rather than require a discrete reaching movement, they 655 were told to continuously adjust their hand position, focusing on accuracy in aligning the hand with the 656 target. The direction of the error was relatively consistent across targets, with the final hand position 657 shifted to the right and undershooting the target. The magnitude of these biases increased as the radial

658 extent of the limb increased. This basic pattern has been observed across studies using different visuo-

659 proprioceptive matching methods^{18,19,27,28,52,56}.

660

The matching errors provide an empirical measure of the transformation from a visual reference frame to a proprioceptive reference frame. To model these data, we defined a transformation error vector, $[x_e, y_e]$, whose direction is fixed across space. We then defined a "reference position" with a coordinate of $[x_r, y_r]$. For upper-limb movements, this reference position is often considered to be positioned around the shoulder.⁵⁷ The transformation error vector at position i is scaled by its Euclidean distance (*d*) to the referent position:

667

$$[7]T_i = d_i[x_e, y_e],$$

668 [8]
$$d_i = \sqrt{(x_i - x_r)^2 + (y_i - x_r)^$$

669 Movements towards a target i is planned via the vector connecting the start position to the target in 670 proprioceptive space, denoted as:

 $(y_r)^2$

671 [9]
$$\sigma h_i = T_0 - (T_i + [x_i, y_i]),$$

where T_0 is the transformation vector at the start position, which is set as [0,0]. Motor bias is calculated as the angular difference between the motor plan and the target. The four free parameters in the Transformation Bias model are x_e , y_e , x_r , y_r .

675

676 Hybrid models

The four models described above each attribute motor biases to a single source. However, the bias might originate from multiple processes. To formalize this hypothesis, we considered three hybrid models, combining the Visual Bias model with the two versions of the Proprioceptive Bias model and with the Transformation Bias model. We did not create a hybrid of the Proprioceptive and Transformation Bias models since they make different assumptions about the information used to derive the motor plan.

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υ	o	L

683 Proprioceptive Bias + Visual Bias (P+V) model

We also created two hybrid models, combining the Visual Bias model with the Vector-Based and Joint-Based Proprioceptive Bias models. The Visual Bias model is used to estimate systematic error in the perceived location of the target and the proprioceptive Bias models are used to estimate systematic error in the perceived position of the hand at the start position. For these models, we calculated the biases

688 from two models separately and then added them together:

689 [10]
$$b_i(P+V) = b_i(\text{Proprioceptive Bias}) + b_i(\text{Visual Bias})$$

- 690 where b_i refers to the bias at target i.
- 691

692 Transformation Bias + Visual Bias (T+V) model

The Transformation Bias model attributes motor biases to error that arises during the transformation of spatial information from visual space to proprioceptive space. However, the perceived location of the visual target may be biased. While we recognize that the misperception of the visual target may influence the transformation bias, the visual bias usually very small in our experiment, with a peak <2°. As such, we simplify by calculating the biases from the Transformation Bias and Visual Bias models separately, and then added the values together:

699

$$[11] b_i(T + V) = b_i(\text{Transformation Bias}) + b_i(\text{Visual Bias})$$

700

701 Model comparison

To compare the models, we fit each model with the data from Experiments 1b and 3b in which reaches were made to 24 targets. We evaluated our models on the average data across participants. We used the fminsearchbnd function in MATLAB to minimize the sum square errors (SSE) and used BIC for model comparison:

706	$[12]$ BIC = $-2LL + k * \ln(n)$
707	where k is the number of parameters of the models, and LL is the sum of loglikelihood across all trials of
708	all participants. Smaller BIC values correspond to better fits.
709	
710	Modeling motor bias after implicit sensorimotor adaptation
711	To examine how the motor bias function changes after visuomotor adaptation, we first used the T+V
712	model to fit the motor bias function from a no-feedback baseline block tested prior to the introduction of
713	the perturbation. We then used the best-fitted baseline model (TVb) to estimate the shift in the motor
714	bias function from data obtained in a no-feedback aftereffect block following adaptation:
715	[13]b(i) = TVb(i+v) + h
716	where $b(i)$ is the motor bias at target i in the aftereffect; v and h indicate the vertical and horizontal shift
717	respectively. To estimate distribution of v and h , we bootstrapped the subjects with repetition for 200
718	times and fitted the v, h based on the group average of each bootstrapped sample.
719	
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726	
727	
728 729	Competing interests
729	RI is a co-founder with equity in Magnetic Tides, Inc.
750	

- 731 **Data availability statement:** Data and code is available at https://github.com/shion707/Motor-Bias
- 732
- 733
- 734 **Reference**
- 1. Burge, J., Ernst, M. O. & Banks, M. S. The statistical determinants of adaptation rate in human
- 736 reaching. J. Vis. 8, 20.1–19 (2008).
- 737 2. Tassinari, H., Hudson, T. E. & Landy, M. S. Combining priors and noisy visual cues in a rapid pointing
 738 task. *J. Neurosci.* 26, 10154–10163 (2006).
- 739 3. Osborne, L. C., Lisberger, S. G. & Bialek, W. A sensory source for motor variation. *Nature* **437**, 412–
- 740 416 (2005).
- Dideriksen, J. L., Negro, F., Enoka, R. M. & Farina, D. Motor unit recruitment strategies and muscle
 properties determine the influence of synaptic noise on force steadiness. *J. Neurophysiol.* 107, 3357–
 3369 (2012).
- Hamilton, A. F. de C., Jones, K. E. & Wolpert, D. M. The scaling of motor noise with muscle strength
 and motor unit number in humans. *Exp. Brain Res.* **157**, 417–430 (2004).
- 746 6. Dhawale, A. K., Smith, M. A. & Ölveczky, B. P. The Role of Variability in Motor Learning. *Annu. Rev.*747 *Neurosci.* 40, 479–498 (2017).
- 748 7. Vindras, P., Desmurget, M., Prablanc, C. & Viviani, P. Pointing Errors Reflect Biases in the Perception
 749 of the InitialHand Position. *J. Neurophysiol.* **79**, 3290–3294 (1998).
- 750 8. Gordon, J., Ghilardi, M. F., Cooper, S. E. & Ghez, C. Accuracy of planar reaching movements. II.
- 751 Systematic extent errors resulting from inertial anisotropy. *Exp. Brain Res.* **99**, 112–130 (1994).
- 9. Ghilardi, M. F., Gordon, J. & Ghez, C. Learning a visuomotor transformation in a local area of work
- 753 space produces directional biases in other areas. *J. Neurophysiol.* **73**, 2535–2539 (06/1995).

- Vindras, P., Desmurget, M. & Viviani, P. Error parsing in visuomotor pointing reveals independent
 processing of amplitude and direction. *J. Neurophysiol.* 94, 1212–1224 (2005).
- Holden, M. P., Newcombe, N. S. & Shipley, T. F. Categorical biases in spatial memory: the role of
 certainty. J. Exp. Psychol. Learn. Mem. Cogn. 41, 473–481 (2015).
- 758 12. Yousif, S. R., Forrence, A. D. & McDougle, S. D. A common format for representing spatial location in
- visual and motor working memory. *Psychon. Bull. Rev.* (2023) doi:10.3758/s13423-023-02366-3.
- Huttenlocher, J., Hedges, L. V., Corrigan, B. & Crawford, L. E. Spatial categories and the estimation of
 location. *Cognition* 93, 75–97 (2004).
- 762 14. Goble, J. A., Zhang, Y., Shimansky, Y., Sharma, S. & Dounskaia, N. V. Directional biases reveal
- 763 utilization of arm's biomechanical properties for optimization of motor behavior. *J. Neurophysiol.* 98,
 764 1240–1252 (2007).
- 765 15. Alexander, R. M. A minimum energy cost hypothesis for human arm trajectories. *Biol. Cybern.* 76,
 766 97–105 (1997).
- 767 16. Nishii, J. & Taniai, Y. Evaluation of trajectory planning models for arm-reaching movements based
 768 on energy cost. *Neural Comput.* 21, 2634–2647 (2009).
- 769 17. Kosovicheva, A. & Whitney, D. Stable individual signatures in object localization. *Curr. Biol.* 27, R700–
 770 R701 (2017).
- 18. Rincon-Gonzalez, L., Buneo, C. A. & Helms Tillery, S. I. The Proprioceptive Map of the Arm Is
 Systematic and Stable, but Idiosyncratic. *PLoS One* 6, e25214- (2011).
- van Beers, R. J., Sittig, A. C. & Denier van der Gon, J. J. The precision of proprioceptive position sense. *Exp. Brain Res.* 122, 367–377 (1998).
- Wang, T. *et al.* Accuracy of hand localization is subject-specific and improved without performance
 feedback. *Sci. Rep.* **10**, 19188 (12/2020).

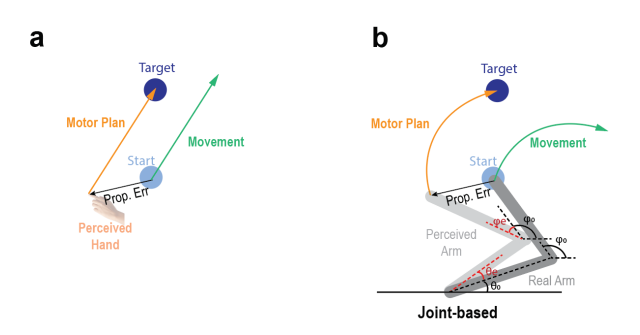
- Sober, S. J. & Sabes, P. N. Flexible strategies for sensory integration during motor planning. *Nat. Neurosci.* 8, 490–497 (2005).
- Sober, S. J. & Sabes, P. N. Multisensory integration during motor planning. *J. Neurosci.* 23, 6982–
 6992 (2003).
- Buneo, C. A., Jarvis, M. R., Batista, A. P. & Andersen, R. A. Direct visuomotor transformations for
 reaching. *Nature* 416, 632–636 (2002).
- 783 24. Soechting, J. F. & Flanders, M. Errors in pointing are due to approximations in sensorimotor
 784 transformations. *J. Neurophysiol.* 62, 595–608 (1989).
- 785 25. Tillery, S. I., Flanders, M. & Soechting, J. F. A coordinate system for the synthesis of visual and
 786 kinesthetic information. *J. Neurosci.* 11, 770–778 (1991).
- 787 26. Flanders, M. & Soechting, J. F. Frames of reference for hand orientation. *J. Cogn. Neurosci.* 7, 182–
 788 195 (1995).
- Jones, S. A. H., Cressman, E. K. & Henriques, D. Y. P. Proprioceptive localization of the left and right
 hands. *Exp. Brain Res.* 204, 373–383 (2009).
- Cressman, E. K. & Henriques, D. Y. P. Reach adaptation and proprioceptive recalibration following
 exposure to misaligned sensory input. *J. Neurophysiol.* **103**, 1888–1895 (2010).
- Palasubramanian, R., Howe, R. D. & Matsuoka, Y. Task performance is prioritized over energy
 reduction. *IEEE Trans. Biomed. Eng.* 56, 1310–1317 (2009).
- 30. Summerside, E. M., Courter, R. J., Shadmehr, R. & Ahmed, A. A. Slowing of movements in healthy
- aging as a rational economic response to an elevated effort landscape. J. Neurosci. (2024)
- 797 doi:10.1523/JNEUROSCI.1596-23.2024.
- 798 31. Soechting, J. F., Buneo, C. A., Herrmann, U. & Flanders, M. Moving effortlessly in three dimensions:
- does Donders' law apply to arm movement? J. Neurosci. 15, 6271–6280 (1995).

- 800 32. Gordon, J., Ghilardi, M. F. & Ghez, C. Impairments of reaching movements in patients without 801 proprioception. I. Spatial errors. *J. Neurophysiol.* **73**. 347–360 (1995).
- 802 33. Slijper, H., Richter, J., Over, E., Smeets, J. & Frens, M. Statistics predict kinematics of hand
 803 movements during everyday activity. *J. Mot. Behav.* 41, 3–9 (2009).
- 804 34. van der Kooij, K., Brenner, E., van Beers, R. J., Schot, W. D. & Smeets, J. B. J. Alignment to natural and
- 805 imposed mismatches between the senses. J. Neurophysiol. **109**, 1890–1899 (2013).
- 806 35. Flash, T. & Hogan, N. The coordination of arm movements: an experimentally confirmed
 807 mathematical model. *J. Neurosci.* 5, 1688–1703 (1985).
- 808 36. Morehead, J. R., Taylor, J. A., Parvin, D. E. & Ivry, R. B. Characteristics of Implicit Sensorimotor
- 809 Adaptation Revealed by Task-irrelevant Clamped Feedback. J. Cogn. Neurosci. 29, 1061–1074 (2017).
- 810 37. Tsay, J. S., Parvin, D. E. & Ivry, R. B. Continuous reports of sensed hand position during sensorimotor
 811 adaptation. *J. Neurophysiol.* 124, 1122–1130 (2020).
- 812 38. Kim, H. E., Morehead, J. R., Parvin, D. E., Moazzezi, R. & Ivry, R. B. Invariant errors reveal limitations
- 813 in motor correction rather than constraints on error sensitivity. *Commun Biol* **1**, 19 (12/2018).
- 814 39. Batista, A. P., Buneo, C. A., Snyder, L. H. & Andersen, R. A. Reach plans in eye-centered coordinates.
 815 Science 285, 257–260 (1999).
- 816 40. Blohm, G., Khan, A. Z. & Crawford, J. D. Spatial Transformations for Eye–Hand Coordination. in
 817 *Encyclopedia of Neuroscience* 203–211 (Elsevier, 2009).
- 41. Henriques, D. Y., Klier, E. M., Smith, M. A., Lowy, D. & Crawford, J. D. Gaze-centered remapping of
- 819 remembered visual space in an open-loop pointing task. *J. Neurosci.* **18**, 1583–1594 (1998).
- Proske, U. & Gandevia, S. C. The proprioceptive senses: their roles in signaling body shape, body
 position and movement, and muscle force. *Physiol. Rev.* 92, 1651–1697 (2012).
- 43. Héroux, M. E., Butler, A. A., Robertson, L. S., Fisher, G. & Gandevia, S. C. Proprioception: a new look
- 823 at an old concept. J. Appl. Physiol. **132**, 811–814 (2022).

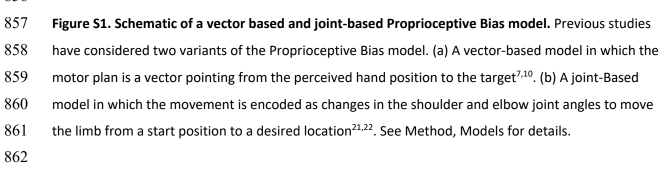
- 44. Zhang, H., Daw, N. D. & Maloney, L. T. Human representation of visuo-motor uncertainty as mixtures
 of orthogonal basis distributions. *Nat. Neurosci.* 18, 1152–1158 (2015).
- 45. van den Berg, R., Shin, H., Chou, W.-C., George, R. & Ma, W. J. Variability in encoding precision
- 827 accounts for visual short-term memory limitations. Proc. Natl. Acad. Sci. U. S. A. 109, 8780–8785
- 828 (2012).
- 46. Hahn, M. & Wei, X.-X. A unifying theory explains seemingly contradictory biases in perceptual estimation. *Nat. Neurosci.* (2024) doi:10.1038/s41593-024-01574-x.
- 47. Mawase, F., Lopez, D., Celnik, P. A. & Haith, A. M. Movement Repetition Facilitates Response
 Preparation. *Cell Rep.* 24, 801–808 (2018).
- 833 48. Shadmehr, R., Huang, H. J. & Ahmed, A. A. A representation of effort in decision-making and motor
 834 control. *Curr. Biol.* 26, 1929–1934 (2016).
- 49. van Beers, R. J., Wolpert, D. M. & Haggard, P. When Feeling Is More Important Than Seeing in
 Sensorimotor Adaptation. *Curr. Biol.* 12, 834–837 (2002).
- Sheehan, T. C. & Serences, J. T. Distinguishing response from stimulus driven history biases. *bioRxiv*(2023) doi:10.1101/2023.01.11.523637.
- Tsay, J. S., Kim, H., Haith, A. M. & Ivry, R. B. Understanding implicit sensorimotor adaptation as a
 process of proprioceptive re-alignment. *Elife* **11**, e76639 (2022).
- S2. Cressman, E. K. & Henriques, D. Y. P. Sensory recalibration of hand position following visuomotor
 adaptation. *J. Neurophysiol.* **102**, 3505–3518 (2009).
- S3. Oldfield, R. C. The assessment and analysis of handedness: The Edinburgh inventory. *Neuropsychologia* 9, 97–113 (1971).
- Schoppe, O., Harper, N. S., Willmore, B. D. B., King, A. J. & Schnupp, J. W. H. Measuring the
 Performance of Neural Models. doi:10.3389/fncom.2016.00010.

- 847 55. Fryar, C. D., Gu, Q. & Ogden, C. L. Anthropometric reference data for children and adults: United
- 848 States, 2007-2010. *Vital Health Stat.* 11 1–48 (2012).
- 849 56. Johnson, E. O., Babis, G. C., Soultanis, K. C. & Soucacos, P. N. Functional neuroanatomy of
- 850 proprioception. J. Surg. Orthop. Adv. **17**, 159–164 (2008).
- 851 57. Haggard, P., Newman, C., Blundell, J. & Andrew, H. The perceived position of the hand in space.
- 852 *Percept. Psychophys.* **62**, 363–377 (2000).









863





865 Figure S2. The Transformation Bias model can explain the motor bias functions when the visual

866 **information is shifted.** (a) In Sober and Sabes (2003)²² participants performed center-out reaches to a

visual target. To perturb the visual information, the start position was presented 6 cm to the left or right

868 of the actual start position of the hand. (b) Participants showed a one-peaked motor bias functions with

the shift-left and shift-right functions shifted in an antiphase relationship to one another. (c) These bias

870 functions are quantitively captured by the Transformation Bias model.

	<i>x_r</i> (cm)	<i>y_r</i> (cm)	x _e	y _e	a (°)	b (°)
Exp 1b	75.7	-17.2	-0.27	-0.05	19.6	3.8
Exp 3b	75.1ª	-0.14 ^{a,b}	-0.24	-0.00	20.0	1.5

872

Table S1. Parameter estimates from best fits of the T+V model for the data from Exps 1b and 3b.

874 See Methods for description of each parameter.

a. Participant moved on the trackpad in Exp 3b. We assumed the movement distance was 1 cm and

- 876 scaled the parameters accordingly.
- b. The estimate of y_r is much smaller in Exp 3b compared to Exp 1b, suggesting the workspace in Exp 3b
- is closer to the body. This attenuates the average magnitude of the bias.