¹ Influence of sensory modality and control ² dynamics on human path integration

3

4 Akis Stavropoulos^{1*}, Kaushik J. Lakshminarasimhan^{2*}, Jean Laurens³, Xaq Pitkow^{4,5}, Dora E. Angelaki^{1,4,6}

- 5 ¹Center for Neural Science, New York University, New York, NY, USA
- 6 ²Center for Theoretical Neuroscience, Columbia University, New York, NY, USA
- 7³ Ernst Strüngmann Institute for Neuroscience, Frankfurt, Germany
- 8 ⁴ Department of Neuroscience, Baylor College of Medicine, Houston, TX, USA
- 9⁵ Department of Electrical and Computer Engineering, Rice University, Houston, TX, USA
- 10 ⁶ Tandon School of Engineering, New York University, New York, NY, USA
- 11 * Equal Author Contribution

12

- 13 Correspondence:
- 14 Prof. Dora E. Angelaki
- 15 Email: da93@nyu.edu
- 16 Center for Neural Science, Mayer 901
- 17 New York University, NY 10003

18 Abstract

Path integration is a sensorimotor computation that can be used to infer latent dynamical states by 19 integrating self-motion cues. We studied the influence of sensory observation (visual/vestibular) and latent 20 control dynamics (velocity/acceleration) on human path integration using a novel motion-cueing 21 algorithm. Sensory modality and control dynamics were both varied randomly across trials, as participants 22 controlled a joystick to steer to a memorized target location in virtual reality. Visual and vestibular steering 23 cues allowed comparable accuracies only when participants controlled their acceleration, suggesting that 24 vestibular signals, on their own, fail to support accurate path integration in the absence of sustained 25 acceleration. Nevertheless, performance in all conditions reflected a failure to fully adapt to changes in 26 the underlying control dynamics, a result that was well explained by a bias in the dynamics estimation. 27 This work demonstrates how an incorrect internal model of control dynamics affects navigation in volatile 28 environments in spite of continuous sensory feedback. 29

30 Introduction

31 Imagine driving a car onto an icy road, where steering dynamics can change rapidly. To avoid crashing,

32 one must rapidly infer the new dynamics and respond appropriately to keep the car on the desired path.

33 Conversely, when you leave an ice patch, control dynamics change again, compelling you to re-adjust

your steering. The quality of sensory cues may also vary depending on environmental factors (e.g. reduced
visibility in fog or twilight, sub-threshold vestibular stimulation under near-constant travel velocity).
Humans are adept at using time-varying sensory cues to adapt quickly to a wide range of latent control
dynamics in volatile environments. However, the relative contributions of different sensory modalities
and the precise impact of latent control dynamics on goal-directed navigation remain poorly understood.
Here we study this in the context of path integration.

Path integration, a natural computation in which the brain uses dynamic sensory cues to infer the evolution 40 of latent world states to continuously maintain a self-position estimate, has been studied in humans, but 41 past experimental paradigms imposed several constraints. First, in many tasks, the motion was passive 42 and/or restricted along predetermined, often one-dimensional, trajectories (Klatzky et al., 1998; Jürgens 43 & Becker, 2006; Petzschner & Glasauer, 2011; Campos et al., 2012; Tramper & Medendorp, 2015). 44 Second, unlike time-varying actions that characterize navigation under natural conditions, participants' 45 responses were often reduced to single, binary end-of-trial decisions (Horst et al., 2015; Chrastil et al., 46 2016; Koppen et al., 2019). Third, even studies that explored contributions of different sensory modalities 47 in naturalistic settings failed to properly disentangle vestibular from motor cues generated during active 48 locomotion (Kearns et al., 2002; Campos et al., 2010; Bergmann et al., 2011; Chen et al., 2017, Arthur et 49 al., 2012; Péruch et al., 1999, 2005). Furthermore, varying constraints have presumably resulted in 50 inconsistent findings on the contribution of vestibular cues to path integration (Jürgens & Becker, 2006; 51 Campos et al., 2010; Horst et al., 2015; Tramper & Medendorp, 2015; Koppen et al., 2019; Chrastil et al., 52 2019; Glasauer et al., 1994; Seidman, 2008). 53

There is a tight link between path integration and spatial navigation on the one hand, and internal models 54 and control dynamics on the other. To accurately estimate self-motion, we rely not only on momentary 55 sensory evidence but also on the knowledge of motion dynamics, *i.e.*, an internal model of the world. 56 Knowledge of the dynamics makes the sensory consequences of actions predictable, allowing for more 57 dexterous steering. However, although there is a large body of research focused on dynamics and 58 adaptation for motor control (Shadmehr & Mussa-Ivaldi, 1994; Lackner & Dizio, 1994; Krakauer et al., 59 1999; Takahashi et al., 2001; Burdet et al., 2001; Kording et al., 2007; Berniker et al., 2010), studies of 60 perceptual inference of latent dynamics during navigation have been limited. Some pioneering studies 61 demonstrated participants' ability to reproduce arbitrary one-dimensional velocity profiles (Grasso et al., 62 1999; Israël et al., 1997), while more recent efforts showed that the history of linear (Petzschner & 63 Glasauer, 2011) and angular (Prsa et al., 2015) displacements affects how participants process sensory 64 input in the current trial. We previously observed that false expectations about the magnitude of self-65 motion can have a drastic effect on path integration (Lakshminarasimhan et al., 2018). We wondered 66 whether prior expectations about the temporal dynamics of self-motion, i.e. how velocities are temporally 67 correlated, can also propagate over time to influence navigation. 68

To explore how dynamics influence navigation across sensory modalities (visual, vestibular, or both), we have built upon a naturalistic paradigm of path integration in which participants navigate to a briefly-cued target location using a joystick to control their velocity in a virtual visual environment (Lakshminarasimhan et al., 2018; Alefantis et al., 2021). Here, we generalize this framework by varying both the control dynamics (joystick control varied along a continuum from velocity to acceleration) and the available sensory cues (vestibular, visual, or both). To achieve this, we designed a motion-cueing

75 algorithm (MC) to render self-motion stimuli according to a joystick control input of maintained accelerations while maintaining correspondence between visual (optic flow) and inertial cues. Using a 76 motion platform with six degrees of freedom to approximate the accelerations that an observer would feel 77 under the imposed control dynamics, we ensured that the MC algorithm would generate matching visual 78 and vestibular cues to closely approximate the desired self-motion (see Methods, Fig. 1 supplement 1 79 and 2). The development of the MC algorithm represents a departure from classical paradigms of 80 navigation research in humans (Chrastil et al., 2019; Israël et al., 1996; Koppen et al., 2019; Seemungal 81 et al., 2007; Horst et al., 2015), as it helps eliminate artificial constraints while still allowing for the 82 isolation of different sensory contributions, most notably vestibular/somatosensory cues, during active, 83 volitional, steering. 84

We found that participants' steering responses were biased (undershooting), and the biases were more prominent in the vestibular condition. Furthermore, steering biases were strongly modulated by the underlying control dynamics. These findings suggest that inertial cues alone (as generated by motion cueing) lack the reliability to support accurate path integration in the absence of sustained acceleration, and that an accurate internal model of control dynamics is needed to make use of sensory observations when navigating in volatile environments.

91 **Results**

92 Task structure

93 Human participants steered towards a briefly-cued target location on a virtual ground plane, with varying sensory conditions and control dynamics interleaved across trials. Participants sat on a motion platform in 94 front of a screen displaying a virtual environment (Fig. 1A). Stereoscopic depth cues were provided using 95 polarizing goggles. On each trial, a circular target appeared briefly at a random location (drawn from a 96 uniform distribution within the field of view; Fig. 1B,C) and participants had to navigate to the 97 remembered target location in the virtual world using a joystick to control linear and angular self-motion. 98 The virtual ground plane was defined visually by a texture of many small triangles which independently 99 appeared only transiently; they could therefore only provide optic-flow information and could not be used 100 as landmarks. The self-motion process evolved according to Markov dynamics, such that the movement 101 velocity at the next time step depended only on the current joystick input and the current velocity 102 103 (Methods – Equation 1).

104 A time constant for the control filter (control timescale) governed the control dynamics: in trials with a small time constant and a fast filter, joystick position essentially controlled velocity, providing participants 105 with responsive control over their self-motion, resembling regular road-driving dynamics. However, when 106 the time constant was large and the control filter was slow, joystick position mainly controlled 107 acceleration, mimicking high inertia under viscous damping, as one would experience on an icy road 108 where steering is sluggish (Fig. 1D right and 1E - top vs bottom). For these experiments, as the control 109 timescale changed, the maximum velocity was adjusted so that the participant could reach the typical 110 target in about the same amount of time on average. This design ensured that the effect of changing control 111 dynamics would not be confused with the effect of integrating sensory signals over a longer or shorter 112 113 time.

bioRxiv preprint doi: https://doi.org/10.1101/2020.09.21.307256; this version posted December 3, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.



Figure 1: (A) Experimental set up. Participants sit on a 6-degrees-of-freedom motion platform with a coupled rotator that allowed unlimited yaw displacements. Visual stimuli were back-projected on a screen (see Methods). The joystick participants used to navigate in the virtual world is mounted in front of the participants' midline. (B) Schematic view of the experimental virtual environment. Participants use a joystick to navigate to a cued target (yellow disc) using optic flow cues generated by ground plane elements (brown triangles; Visual and Combined conditions only). The ground plane elements appeared transiently at random orientations to ensure they cannot serve as spatial or angular landmarks. (C) Left: Overhead view of the spatial distribution of target positions across trials. *Red dot* shows the starting position of the participant. Positions were uniformly distributed within the participant's field of view. Right: Movement trajectories of one participant during a representative subset of trials. Starting location is denoted by the red dot. (D) Control dynamics. Inset: Linear joystick input from a subset of trials in the visual condition of an example participant. Left: Simulated maximum pulse joystick input (max joystick input = I) (see also Fig. 1 supplement 3). This input is lowpass filtered to mimic the existence of inertia. The time constant of the filter varies across trials (time constant τ). In our framework, maximum velocity also varies according to the time constant τ of each trial to ensure comparable travel times across trials (see Methods - Control Dynamics). Right: the same joystick input (scaled by the corresponding maximum velocity for each τ) produces different velocity profiles for different time constants ($\tau = 0.6s$) corresponds to velocity control; $\tau = 3s$ corresponds to acceleration control; τ values varied randomly along a continuum across trials, see **Methods**). Also depicted is the brief cueing period of the target at the beginning of the trial (gray zone, *1 second*) long). (E) Markov decision process governing self-motion sensation (Methods – Equation 1), u, v, and o denote joystick input, movement velocity, and sensory observations, respectively, and subscripts denote time indices. Note that due to the 2-D nature of the task, these variables are all vector-valued, but we depict them as scalars for the purpose of illustration. By varying the time constant, we manipulated the control dynamics (*i.e.*, the degree to which the current velocity carried over to the future, indicated by the thickness of the horizontal lines) along a continuum such that the joystick position primarily determined either the participant's velocity (top; thin lines) or acceleration (bottom; thick lines) (compare with (D) top and bottom, respectively). Sensory observations were available in the form of vestibular (left), optic flow (middle), or both (right).

114 Concurrently, we manipulated the modality of sensory observations to generate three conditions: 1) a 115 *vestibular* condition in which participants navigated in darkness, and sensed only the platform's motion 116 (note that this condition also engages somatosensory cues, see **Methods**), 2) a *visual* condition in which 117 the motion platform was stationary and velocity was signaled by optic flow, and 3) a *combined* condition 118 in which both cues were available (**Fig. 1E** – left to right). Across trials, sensory conditions were randomly

119 interleaved while manipulation of the time constant followed a bounded random walk (**Methods** – 120 Equation 2). Participants did not receive any performance-related feedback.





Figure 2: (A) Geometric definition of analysis variables. The *gray solid line* indicates an example trajectory. The target and response distance and angle relative to the starting position of the participant are given by r, θ (*thin lines*) and \tilde{r} , $\tilde{\theta}$ (*thick lines*), respectively. (B), (C) Example participant: Comparison of the radial distance \tilde{r} of an example participant's response (final position) against the radial distance r of the target (B), as well as the angular eccentricity of the participant's response $\tilde{\theta}$ versus target angle θ (C), across all trials for one participant, colored according to the sensory condition (*green:* vestibular, *cyan:* visual, *purple:* combined visual and vestibular; Figure 2 source data 1). Radial and angular response gains were defined as the slope of the corresponding regressions. Black dashed lines show unity slope, and the solid lines represent slopes of the regression fits (intercept set to 0). (D) All participants: Radial and angular gains in each sensory condition of data points for the corresponding sensory condition. *Diamonds* (centers of the ellipses) represent the mean radial and angular response gains across participants. Dashed lines indicate unbiased radial or angular position responses. Solid diagonal line has unit slope. (E) Magnitudes of radial and angular components of control inputs across sensory conditions for an example participant. *Shaded regions* represent ± 1 standard deviation across trials. The gray zone corresponds to the target presentation period.

We first compared the participants' stopping locations on each trial to the corresponding target locations, separately for each sensory condition. We calculated the radial distance \tilde{r} and angular eccentricity $\tilde{\theta}$ of

124 the participants' final position relative to the initial position (Fig. 2A), and compared them to the initial target distance r and angle θ , as shown for all trials (all time constants together) of a typical participant in 125 Fig. 2B, C. This revealed biased performance with notable undershooting (participants stopped short of 126 the true target location), in both distance and angle, which was well described by a linear model without 127 intercept (Radial Distance $R^2 \pm$ standard deviation – vestibular: 0.39±0.06, visual: 0.67±0.1, combined: 128 0.64±0.11; Angular Eccentricity R^2 ± standard deviation – vestibular: 0.85±0.06, visual: 0.95±0.05, 129 combined: 0.96±0.04. Adding a non-zero intercept term offered negligible improvement: Radial Distance 130 ΔR^2 – vestibular: 0.02±0.02, visual: 0.03±0.03, combined: 0.03±0.02; Angular Eccentricity ΔR^2 – 131 vestibular: 0.02±0.03, visual: 0.01±0.01, combined: 0.01±0.01). We refer to the slope of the linear 132 regression as 'response gain': a response gain of unity indicates no bias, while gains larger (smaller) than 133 unity indicate overshooting (undershooting). As shown with the example participant in Fig. 2B, C, there 134 was substantial undershooting in the vestibular condition, whereas performance was relatively unbiased 135 under the combined and visual conditions (see also Fig. 2 supplement 1A). These results were consistent 136 across participants (Fig. 2D, mean radial gain \pm standard deviation – vestibular: 0.76 \pm 0.25, visual: 137 0.88 ± 0.23 , combined: 0.85 ± 0.22 , mean angular gain \pm standard deviation – vestibular: 0.79 ± 0.22 , visual: 138 0.98±0.14, combined: 0.95±0.12), and no significant sex differences were observed (see Fig. 2 139 supplement 1B). The difference in response gain between modalities could be traced back to the control 140 exerted by the subjects on the joystick. Both linear and angular components of control input had shorter 141 duration in the vestibular condition (mean \pm SEM of total area of joystick input across participants (a.u.): 142 Radial – vestibular: 5.62 ± 0.27 , visual: 7.31 ± 0.33 , combined: 7.07 ± 0.34 ; Angular – vestibular: 2.39 ± 0.30 , 143 visual: 3.29 ± 0.42 , combined: 3.79 ± 0.46), and produced smaller displacements, as summarized by the 144 response gains (Fig. 2E, Fig. 2 supplement 2). 145

146 Effect of control dynamics on performance

To examine whether control dynamics affected the response gain, we performed three complementary 147 analyses. First, we recomputed response gains by stratifying the trials into three groups of equal size based 148 on the time constants. We found that smaller time constants (velocity control) were associated with smaller 149 response gains (Fig. 3A; Table 1). This relationship was most pronounced in the vestibular condition, 150 where larger time constants (acceleration control) resulted in better (closer to ideal) performance (Fig. 3A, 151 green; see Discussion). Control dynamics had a smaller but considerable effect on steering responses in 152 the visual and combined conditions, with participants exhibiting modest overshooting (undershooting) 153 when the time constant was large (small) (Fig. 3A, cyan/purple). 154

Second, we performed a fine-grained version of the above analysis by computing residual errors on each 155 trial, *i.e.* the deviation of the response from the mean response predicted from target location alone 156 (Methods – Equation 3). Since participants try to stop at their believed target location, ideally their mean 157 responses should depend only on target location, and not on control dynamics. In other words, if 158 participants adapted their control appropriately to the varying control dynamics, their responses should 159 cluster around their mean response, and as a result, their residual errors should be centered around zero 160 without any mean dependence on dynamics. However, we found a significant correlation between residual 161 errors and the time constant across trials (Fig. 3B and C, Fig. 3 supplement 1, Table 2, see Methods; no 162 significant sex differences were observed, and therefore are not investigated in subsequent analyses, see 163 also Fig. 2 supplement 1C). This correlation, and the corresponding regression slopes, were substantially 164



Figure 3: Effect of control dynamics on participants' responses. (A) Participant average of radial and angular response gains in each condition, with trials grouped into tertiles of increasing time constant τ . Error bars denote ± 1 SEM. (B) Effect of time constant τ on radial (left) and angular (right) residual error, for an example participant (Figure 3 source data 1). Solid lines represent linear regression fits and ellipses the 68% confidence interval of the distribution for each sensory condition. Dashed lines denote zero residual error (i.e. stopping location matches mean response). (C) Correlations of radial (ε_r) and angular (ε_{θ}) residual errors with the time constant for all participants. Ellipses indicate the 68% confidence intervals of the distribution of data points for each sensory condition. Solid diagonal line has unit slope. Across participants, radial correlations, which were larger for the vestibular condition, were greater than angular correlations (See also Table 2). (D) Linear regression coefficients for the prediction of participants' response location (final position: $\tilde{r}, \tilde{\theta}$; left and right, respectively) from initial target location (r, θ) and the interaction between initial target location and the time constant $(r\tau, \theta\tau)$ (all variables were standardized before regressing, see Methods; Figure 3 source data 2). Asterisks denote statistical significance of the difference in coefficient values of the interaction terms across sensory conditions (paired t-test; *: p<0.05, **: p<0.01, ***: p<0.001; see main text). Error bars denote ± 1 SEM. Note a qualitative agreement between the terms that included target location only and the gains calculated with the simple linear regression model (Fig. 2B). (E) Comparison of actual and null-case (no adaptation) response gains, for radial (top) and angular (bottom) components, respectively (average across participants). Dashed lines represent unity lines, i.e. actual response gain corresponds to no adaptation. Inset: Regression slopes between actual and null-case response gains. A slope of 0 or 1 corresponds to perfect or no adaptation (gray dashed lines), respectively. *Error bars* denote ± 1 SEM.

higher in the vestibular condition (Mean Pearson's $r \pm$ SEM: Radial component – vestibular: 0.52±0.02, visual: 0.36±0.03, combined: 0.37±0.03; Angular component – vestibular: 0.23±0.02, visual: 0.23±0.03, combined: 0.26±0.03; see also **Table 2,3**). Thus, for a given target distance, participants tended to travel

168 further when the time constant was larger (acceleration control), indicating they did not fully adapt their 169 steering control to the underlying dynamics.

Third, to quantify the contribution of the time constant in the participants' responses, we expanded the 170 linear model to accommodate a dependence of response (final stopping position) on target location, time 171 constant, and their interaction. A partial correlation analyses revealed that the time constant contributed 172 substantially to participants' response gain, albeit only by modulating the radial and angular distance 173 dependence (Table 4; Fig. 3 supplement 2; see Methods – Equation 4). Again, the contribution of the 174 time constant-dependent term was much greater for the vestibular condition (Fig. 3D), especially for the 175 radial distance (p-values of difference in coefficient values across modalities obtained by a paired t-test – 176 Radial: vestibular vs visual: $p < 10^{-4}$, vestibular vs combined: $p < 10^{-4}$; Angular: vestibular vs visual: 177 p = 0.016, vestibular vs combined: p = 0.013). While perfect adaptation should lead to response gain 178 that is independent of control dynamics, all three independent analyses revealed that control dynamics did 179 substantially influence the steering response gain, exposing participants' failure to adapt their steering to 180 the underlying dynamics. Adaptation was lowest for the vestibular condition; in contrast, for the visual 181 and combined conditions, the response gain was less affected indicating greater compensation when visual 182 information was available. 183

We quantified the extent to which participants failed to adapt to the control dynamics, by simulating a null 184 case for no adaptation. Specifically, we generated null-case trajectories by using the steering input from 185 actual trials and re-integrating it with time constants from other trials. In this set of null-case trajectories, 186 the steering control corresponds to different time constants; in other words, steering is not adapted to the 187 underlying dynamics (see Methods). We then grouped these trajectories based on the simulation time 188 constant (as in **Fig. 3A**) and computed the corresponding response gains. We found that the true response 189 gains in the vestibular condition where much closer to the no-adaptation null case, compared to 190 visual/combined conditions (Fig. 3E). Interestingly, this finding was more prominent in the radial 191 component of the response gain (Fig. 3E insets), consistent with our earlier observations of a stronger 192 influence of the dynamics on the radial component of the responses. 193

We have shown how various measures of the participants' final responses (stopping positions, response gain, residual errors) are influenced by the time constant of the dynamics. This large dependence of the final responses on the time constant exposes participants' failure to fully adapt their steering to the underlying dynamics. In other words, the influence of the dynamics on steering control was relatively weak, especially in the vestibular condition.

For best performance, however, control dynamics *should* influence the time course of steering behavior. 199 We directly quantified the influence of the control dynamics on steering by comparing participants' 200 braking (negative control input) across time constants: when the time constant is large, we ideally expect 201 to see more braking as a countermeasure for the sluggish control (Fig. 1D) to minimize travel duration 202 (see Methods). Indeed, participants do tend to brake more for higher time constants, but this effect is 203 weaker in the vestibular condition (Fig. 4 and 4 inset). Nevertheless, correlations between the time 204 constant and cumulative braking (total area below zero linear control input) were significant in all sensory 205 conditions (Mean Pearson's $r \pm$ SEM – vestibular: 0.20±0.03, visual: 0.62±0.04, combined: 0.57±0.04; p-206 values of Pearson's r difference from zero – vestibular: $p=10^{-5}$, visual: $p<10^{-7}$, combined: $p<10^{-7}$). Overall, 207 208 it appears that behavior in the vestibular condition is minimally influenced by the dynamics (i.e. smaller 209 modulation of control input by the time constant, as shown by the cumulative braking). When optic flow 210 is available, however, participants are more flexible in adjusting their control.



211 We have shown previously that accumulating sensory noise over an extended time (~ 10 s) would lead to

a large uncertainty in the participant's beliefs about their position, causing them to undershoot 212 (Lakshminarasimhan et al. 2018). The exact amount of undershooting depends both on the reliability of 213 self-motion cues, which determines the *instantaneous* uncertainty in the self-motion estimate, and on 214 travel duration, which governs how much uncertainty is accumulated while navigating to the target. With 215 recent findings ascribing uncertainty accumulation to noise in the velocity input (Stangl et al., 2020), the 216 observed differences in navigation performance across sensory modalities can be readily attributed to 217 greater measurement noise (lower reliability) in vestibular signals. On the other hand, we observed 218 performance differences across control dynamics within each sensory modality, so those differences 219 cannot be attributed to differences in the reliability of self-motion cues (instantaneous uncertainty). 220 However, it might seem that this effect of control dynamics must be due to either differences in travel 221 duration or velocity profiles, which would both affect the accumulated uncertainty. We adjusted stimulus 222 parameters to ensure that the average travel time and average velocity were similar across different control 223 dynamics (Methods – Equation 1.2-1.10), however, we found that travel duration and average velocity 224 depend weakly on the time constant in some participants. Simulations suggest that both dependencies are 225 a consequence of maladaptation to the dynamics rather than a cause of the observed effect of the dynamics 226 on the responses. Interestingly, the dependence is stronger in the vestibular condition where there is less 227 adaptation to the dynamics, agreeing with our simulations (Fig. 5 supplement 1A,B). Differences in 228 velocity profiles is also an unlikely explanation since their expected effect on the participants' responses 229 (undershoot) is the opposite of the observed effect of the control dynamics (overshooting tendency; Fig. 230 5 supplement 1C). Consequently, unlike the effect of sensory modality on response gain, neither 231 instantaneous nor accumulated differences in the uncertainty can fully account for the influence of control 232

dynamics, *i.e.* the time constant. Instead, we will now show that the data are well explained by strong prior expectations about motion dynamics that cause a bias in estimating the time constant.

235 Modelling the effect of control dynamics across sensory modalities

236 From a normative standpoint, to optimally infer movement velocity, one must combine sensory observations with the knowledge of the time constant. Misestimating the time constant would produce 237 errors in velocity estimates, which would then propagate to position estimates, leading control dynamics 238 to influence response gain (Fig. 5A, middle-right). This is akin to misestimating the slipperiness of an ice 239 patch on the road causing an inappropriate steering response, that would culminate in a displacement that 240 differs from the intended one (Fig. 5 supplement 2). However, in the absence of performance-related 241 feedback at the end of the trial, participants would be unaware of this discrepancy, wrongly believing that 242 the actual trajectory was indeed the intended one. In other words, participants' imperfect adaptation to 243 changes in control dynamics could be a consequence of control dynamics misestimation. 244

We tested the hypothesis that participants misestimated the time constant using a two-step model that 245 reconstructs the participants' believed trajectories according to their *point estimate* of the time constant τ , 246 as follows. First, a Bayesian observer model infers the participant's belief about τ on individual trials, *i.e.* 247 the subjective posterior distribution over the time constant (τ inference step; Fig. 5A, left). Second, we 248 used the median of that belief to reconstruct the believed trajectory by integrating the actual joystick input 249 according to the *estimated* time constant on that trial (integration step), resulting in a believed stopping 250 location (Fig. 5A, middle-right). In the absence of bias (response gain of one), the believed stopping 251 locations should land on or near the target. However, various unmeasurable fluctuations in that belief 252 across trials should lead to variability clustered around the target location. When the behavior is biased 253 (response gain different from one, as was the case here - Fig. 2D), this cluster should instead be centered 254 around the participants' mean belief for that target location (determined from their biased responses and 255 henceforth referred to as mean stopping location). Since the participants' goal is to stop as close to their 256 perceived target location as possible, the deviation of believed stopping locations from the mean stopping 257 location for a given target should be small. We call this deviation the subjective residual error. Therefore, 258 we inferred the parameters of the Bayesian model separately for each participant by minimizing the 259 subjective residual errors induced by the control dynamics using the principle of least-squares (see 260 Methods for further details). We next describe the parameters of the Bayesian model and then describe 261 the results of fitting the model to our data. 262

263 Because the time constant τ is always positive, we model both the prior distribution and the likelihood function over the variable $\varphi = \log \tau$ as Gaussians in log-space. We parameterized both the prior and the 264 likelihood with a mean (μ) and standard deviation (σ). The mean of the prior (μ) was allowed to freely 265 vary across sensory conditions but assumed to remain fixed across trials. On each trial, the likelihood was 266 assumed to be centered on the actual value of the log time-constant τ^* on that trial according to $\mu = \varphi^* =$ 267 $\log \tau^*$ and was therefore not a free parameter. Finally, we set the ratio λ of prior over likelihood σ , to 268 freely vary across sensory conditions. Thus, for each sensory condition, we fit two parameters: the μ of 269 the prior, and the ratio (λ) of prior σ to likelihood σ . As mentioned above, we fit the model to minimize 270 the difference between their believed stopping locations and their experimentally-measured mean stopping 271 272 location (subjective residual errors), using a least-squares approach (Methods) and obtained one set of



Figure 5: (A) Left: Illustration of the Bayesian estimator model. We fit two parameters: the ratio λ of standard deviations of prior and likelihood ($\lambda = \sigma_p / \sigma_l$) and the mean of the prior (μ_{prior}) of the normally distributed variable $\varphi = \log \tau$ (black *dotted box*). Likelihood function is centered on the log-transformation of the actual τ , $\varphi^* = \log \tau^*$ (*Black dashed line*). The time constant estimate $\hat{\tau}$ corresponded to the median of the posterior distribution over τ , which corresponds to the median $\hat{\varphi}$ over φ , $\hat{\tau} = \exp(\hat{\varphi})$, (red dotted box; red dashed line; see **Methods**). *Middle*: Control dynamics implied by the actual time constant τ (top; gray shade) and the estimated time constant $\hat{\tau}$ (bottom; red shade). u, v, and o denote joystick input, movement velocity, and sensory observations respectively, and subscripts denote time indices. \hat{v} denotes the inferred velocity implied by the model. Misestimation of the time constant leads to erroneous velocity estimates about self-motion \hat{v} which result in biased position beliefs. *Right*: Illustration of the actual (black) and believed (red) trajectories produced by integrating (box) the actual velocity v and the estimated velocity \hat{v} , respectively. White and yellow dots denote the starting and target position, respectively. Inset: Illustration of correlated (black dots) and uncorrelated (red dots) residual errors with the time constant for actual and model-implied responses (simulated data). For simplicity, we depict residual errors as onedimensional and assume unbiased responses (response gain of 1). Blown-up dots with yellow halo correspond to the actual and model-implied trajectories of the right panel. Solid black horizontal line corresponds to zero residual error (i.e. stop on target location). (B) Comparison of correlations between real and subjective residual errors with τ (Figure 5 source data 1). On the right, participant averages of these correlations are shown. Colored bars: 'Subjective' correlations, Open bars: Actual correlations. Error bars denote ± 1 SEM across participants. Asterisks denote the level of statistical significance of differences between real and subjective correlations (*: p<0.05, **: p<0.01, ***: p<0.001).

273 parameters for each condition. Finally, the participant's estimated time-constant $\hat{\tau}$ on each trial was taken 274 to be the median of the best-fit model, which equals the median of the distribution over φ (Fig. 5A, left).

By integrating the subject's joystick inputs on each trial using $\hat{\tau}$ rather than the actual time-constant τ , we computed the believed stopping location and the subjective residual errors implied by the best-fit model.

We then compared the correlations between the time constant and the residual errors for real responses 277 278 (from data in Fig. 3B,C) or subjective responses (from model), separately for radial and angular components. Because participants try to stop at their believed target location, the believed stopping 279 position should depend only on target location and not on the control dynamics. Any departure would 280 suggest that participants knowingly failed to account for the effect of the control dynamics, which would 281 manifest as a dependence of the *subjective* residual errors on the time constant τ . In other words, a good 282 model of the participants' beliefs would predict that the *subjective* residual errors should be uncorrelated 283 with the time constant τ (Fig. 5A inset - red) even if the *real* residual errors are correlated with the time 284 constant (Fig. 5A inset - black). In all cases, we observed that the correlation between residual error and 285 time constant was indeed significantly smaller when these errors were computed using the *subjective* 286 (believed) rather than real stopping location (Fig. 5B). In fact, subjective residual errors were completely 287 uncorrelated with the time constant suggesting that the Bayesian model is a good model of participants' 288 beliefs, and that the apparent influence of control dynamics on behavioral performance was entirely 289 because participants misestimated the time constant of the underlying dynamics. 290

We next examined the model posterior estimates to assess how subjects' internal estimate of the control dynamics departed from the true dynamics. The relationship between real and model-estimated time constants for all participants can be seen in **Fig. 6A**. In the vestibular condition, all participants



Figure 6: Model parameters. (A) Relationship between the model-estimated and actual time constant across all participants in vestibular (green), visual (cyan) and combined (purple) conditions. Participant averages are superimposed (*thick lines*). *Dashed line:* unbiased estimation (**Figure 6 source data 1**). (**B**) Fitted model parameters: ratio λ of prior (σ_p) over likelihood (σ_l) standard deviation and mean (μ) of prior. Error bars denote ±1 SEM. *Dashed lines* represent the corresponding values of the sampling distribution of $\varphi = \log \tau$, which is normal (see **Methods; Figure 6 source data 2**). The prior distribution's μ was comparable in the vestibular condition to the μ of the actual sampling distribution (sampling distribution μ : 0.58 log *s* – *p*-value of prior μ difference obtained by bootstrapping – vestibular: p = 0.014, visual: $p = < 10^{-7}$; combined: $p < 10^{-7}$). *Asterisks* denote the level of statistical significance of differences in the fitted parameters across conditions (*: p < 0.05, **: p < 0.01, ***: p < 0.001).

consistently misestimated τ , exhibiting a substantial regression towards the mean (**Fig. 6A**, green). This effect was much weaker in the visual condition. Only a few participants showed relatively flat estimates, with the majority showing smaller departures from ideal estimates (*dashed line*). The data for the combined condition followed a similar trend, with properties between those in the visual and vestibular conditions (**Fig. 6A**, purple). These results suggest that the better control adaptation in the visual and combined conditions shown in **Fig. 3** is due to participants' improved estimates of the time constant when optic flow was available.

The source of inaccuracies in the estimated time constant can be understood by examining the model 301 parameters (Fig. 6B). The ratio λ of prior over likelihood standard deviations was significantly lower in 302 the vestibular condition than other conditions, suggesting stronger relative weighting of the prior over the 303 likelihood (Fig. 6B left, green bar; mean ratio $\lambda \pm$ standard SEM – vestibular: 0.30±0.09, visual: 304 1.02±0.17, combined: 0.80±0.10; p-value of ratio λ paired differences obtained by bootstrapping -305 vestibular vs visual: p = 0.0007, vestibular vs combined: p = 0.0087; visual vs combined: p = 0.016). 306 Notably, the ratio was close to 1 for the visual and combined conditions, suggesting equal weighting of 307 prior and likelihood. Thus, participants' estimate of the control dynamics in the vestibular condition was 308 plagued by a combination of strong prior and weak likelihood, which explains the much stronger 309 regression towards the mean in Fig. 6A. 310

311 Alternative models

312 To test whether our assumption of a static prior distribution over time constants was reasonable, we fit an alternative Bayesian model in which the prior distribution was updated iteratively on every trial, as a 313 weighted average of the prior on the previous trial and the current likelihood over φ (Dynamic prior model; 314 see Methods). For this version, the initial prior μ was taken to be the time constant on the first trial, and 315 we once again modeled the likelihood and prior as normal distributions over the log-transformed variable. 316 φ , where the likelihood was centered on the actual φ and was therefore not a free parameter. Thus, we fit 317 one parameter: the ratio λ of prior σ over likelihood σ . On each trial, the relative weighting of prior and 318 likelihood responsible for the update of the prior depended solely on λ ; that is, the relationship between 319 their corresponding σ (*i.e.* relative widths). The performance of the static and dynamic prior models was 320 comparable in all conditions, for both distance and angle, suggesting that a static prior is adequate in 321 explaining the participants' behavior on this task (Fig. 7; light versus dark bars). In line with our 322 expectations, when updating the prior in the dynamic model, the weighting of the previous-trial prior 323 received significantly more weight in the vestibular condition (in the range of [0,1]; mean prior weights \pm 324 SEM - vestibular: 0.93±0.03, visual: 0.48±0.10, combined: 0.61±0.09; p-value of paired weight 325 differences obtained by bootstrapping - vestibular vs visual: $p = 10^{-5}$, vestibular vs combined: p = 4. 326 10^{-4} ; visual vs combined: p = 0.08). The comparable goodness of models with static and dynamic priors 327 suggest that sensory observations were not too reliable to cause rapid changes in prior expectations during 328 the course of the experiment. 329

At the other extreme, to test whether participants used sensory observations at all to estimate control dynamics, we compared the static prior Bayesian model to a parsimonious model that assumed a fixed time constant across all trials (*i.e.* completely ignoring changes in control dynamics). This latter model can be understood as a Bayesian model instantiated with a very strong static prior. In line with our

334 expectations (see Fig. 6A), this latter model performed comparably in the vestibular condition, but 335 substantially worse in the visual and combined conditions (Fig. 7).



336 Due to the correlated nature of the random walk process dictating the evolution of time constants, an

Figure 7: Comparison of the correlations between the actual τ and the subjective residual errors implied by three different τ estimation models (Bayesian estimation with a Static prior (S), Bayesian estimation with a Dynamic prior (D), Fixed estimate (F)). We tested the hypotheses that either the prior distribution should not be static or that the participants ignored changes in the control dynamics and navigated according to a fixed time constant across all trials (fixed τ -estimate model; see **Methods**). For this, we compared the correlations between the subjective residual error and the actual trial τ that each model produces. The Dynamic prior model performs similarly to the Static prior model in all conditions, indicating that a static prior is adequate in explaining our data (p-values of paired t-test between correlation coefficients of the two models: Distance – vestibular: p =0.96, visual: p = 0.19, combined: p = 0.91; Angle – vestibular: p = 0.87, visual: p = 0.09, combined: p = 0.59). For visual and combined conditions, the fixed τ model not only fails to minimize the correlations but, in fact, strongly reverses it, for both distance (*left*) and angle (*right*). Since these correlations arise from the believed trajectories that the fixed τ model produces, this suggests that participants knowingly stop before their believed target location for higher time constants. Model performance was only comparable in the vestibular condition, where the average correlation of the fixed τ model (F) was contained within the 95% confidence intervals (CI) of the static prior Bayesian model (S), for both distance and angle (Distance - F: mean Pearson's correlation coefficient $\rho = 0.03$, S: 95% CI of Pearson's correlation coefficient $\rho = [-0.10 \ 0.25]$; Angle -F: mean Pearson's correlation coefficient $\rho = -0.01$, S: 95% CI of Pearson's correlation coefficient $\rho = [-0.12 \ 0.15]$). Error bars denote ± 1 SEM.

337 alternative by which participants could get away without estimating the time constant in the vestibular condition would be to carry over their estimate from the previous combined/visual trial to the current 338 vestibular trial. To test this, we considered two models: the time constant estimate in the current vestibular 339 trial was taken to be either the real time constant, or the posterior estimate of the time constant from the 340 previous visual/combined trial. Neither model, however, could account for the participants' behavior, as 341 they could not fully explain away the correlation between the residual errors and the time constant (Fig. 342 7 supplement 1). Intuitively, although choosing actions with respect to the previous trial's time constant 343 should result in estimates that regress towards the mean, the predicted effect is weaker than that observed 344 in the data. 345

Finally, we tested a variation of previously suggested sensory feedback control models (Glasauer et al., 2007; Grasso et al., 1999) where a controller relies solely on sensory inputs to adjust their control without explicitly estimating the latent variables governing the control dynamics. Specifically, the model assumes that participants implement a type of bang-bang control that switches at a certain distance from the target

350 (or more accurately, the mean response). However, this model predicts a much stronger dependence of 351 the responses on the dynamics compared to our data, and characteristics of the predicted control input 352 differ significantly from the actual control (**Fig. 7 supplement 2**). Overall, our results suggest that optic 353 flow, but not vestibular signals, primarily contributes to inferring the latent velocity dynamics.

354 **Discussion**

We showed that human participants can navigate using different sensory cues and that changes in the 355 control dynamics affect their performance. Specifically, we showed that participants can path integrate to 356 steer towards a remembered target location quite accurately in the presence of optic flow. In contrast, 357 inertial (vestibular/somatosensory) cues generated by motion cueing alone lacked the reliability to support 358 accurate path integration, leading to substantially biased responses under velocity control. Performance 359 was also influenced by the changing control dynamics in all sensory conditions. Because control dynamics 360 were varied on a trial-by-trial basis, sensory cues were crucial for inferring those dynamics. We used 361 probabilistic inference models to show that the observed responses are consistent with estimates of the 362 control dynamics that were biased toward the center of the experimental distribution. This was particularly 363 strong under the vestibular condition such that the response gain substantially increased as the motion 364 dynamics tended towards acceleration control. Although control dynamics were correlated across trials, 365 our models showed that participants did not take advantage of those correlations to improve their 366 estimates. 367

368 **Relation to past work**

In the paradigm used here, participants actively controlled linear and angular motion, allowing us to study 369 multisensory path integration in two dimensions with few constraints. This paradigm was made possible 370 by the development of a motion-cueing (MC) algorithm to render visual and vestibular cues either 371 synchronously or separately. In contrast, previous studies on human path integration used restricted 372 paradigms in which motion was either one dimensional or passively rendered, and participants' decisions 373 where typically reduced to end-of-trial binary evaluations of relative displacement (Campos et al., 2012; 374 Chrastil et al., 2016, 2019; Jürgens & Becker, 2006; Koppen et al., 2019; Horst et al., 2015; Tramper & 375 Medendorp, 2015). As a result, findings from past studies that evaluate the contributions of different 376 sensory modalities to self-motion perception (Chrastil et al., 2019; Israël et al., 1996; Koppen et al., 2019; 377 Seemungal et al., 2007; Horst et al., 2015) may be more limited in generalizing to real-world navigation. 378

Our results show that, at least in humans, navigation is driven primarily by visual cues under conditions 379 of near-constant travel velocity (velocity control). This dominance of vision suggests that the reliability 380 of the visual cues is much higher than vestibular cues (as generated by our platform), as corroborated by 381 the data from the combined condition in which performance resembles the visual condition. This makes 382 sense because the vestibular system is mainly sensitive to acceleration, exhibiting higher sensitivity to 383 higher-frequency motion compared to the visual system (Karmali et al., 2014). Consequently, it may only 384 be reliable when motion is dominated by acceleration. This interpretation is further supported by the 385 observation that participants' vestibular performance was a lot less biased in the regime of acceleration 386 joystick control, where accelerations are prolonged during navigation. 387

388 Experimental constraints in past navigation studies have also precluded examining the influence of control 389 dynamics. In fact, the importance of accurately inferring control dynamics, which are critical for

390 predicting the sensory consequences of actions, has largely been studied in the context of limb control and 391 motor adaptation (Burdet et al., 2001; Kording et al., 2007; Krakauer et al., 1999; Lackner & Dizio, 1994; Shadmehr & Mussa-Ivaldi, 1994; Takahashi et al., 2001). Here, we provide evidence for the importance 392 of accurately inferring control dynamics in the context of path integration and spatial navigation. Although 393 participants were not instructed to expect changes in the latent dynamics and received no feedback, we 394 showed that they nevertheless partly adapted to those dynamics while exhibiting a bias toward prior 395 expectations about these dynamics. This biased estimation of control dynamics led to biased path 396 integration performance. This result is analogous to findings about the effect of changing control dynamics 397 in motor control: first, adaptation to the dynamics happens even in the absence of performance-related 398 feedback (Batcho et al., 2016; Lackner & Dizio, 1994) and, second, this adaptation relies on prior 399 experience (Arce et al., 2009) and leads to systematic errors when knowledge of the dynamics is inaccurate 400 (Körding et al., 2004). Thus, participants try to exploit the additional information that the dynamics 401 contain about their self-motion in order to achieve the desired displacement. 402

A Bayesian estimator with a static prior over the dynamics sufficiently explained participants' beliefs in 403 our data, while results were comparable with a dynamic prior that was updated at every trial. This could 404 be attributed to the structure of the random walk of the control dynamics across trials, as a static prior is 405 not as computationally demanding and potentially more suitable for fast changes in the time constant. 406 These Bayesian models attempt to explain behavior in an optimal way given the task structure. Meanwhile, 407 alternative suboptimal models (fixed estimate, carry-over estimate, sensory feedback model) failed to 408 explain behavior successfully, especially when optic flow was available. These results strongly favor 409 underlying computations within the context of optimality in the presence of optic flow. 410

Task performance was substantially worse in the vestibular condition, in a manner suggesting that 411 vestibular inputs from motion cueing lack the reliability to precisely estimate control dynamics on 412 individual trials. Nevertheless, the vestibular system could still facilitate inference by integrating trial 413 history to build expectations about their statistics. Consistent with this, the mean of the prior distribution 414 over the dynamics fit to data was very close to the mean of the true sampled distribution, suggesting that 415 even if within-trial vestibular observations are not sufficient, participants possibly combine information 416 about the dynamics across trials to construct their prior beliefs. This is consistent with the findings of Prsa 417 et al., 2015, where vestibular cues were used to infer an underlying pattern of magnitude of motion across 418 trials. However, the measurement of the dynamics in that study substantially differs from ours: here, 419 motion dynamics are inferred using self-motion cues within each trial whereas in (Prsa et al., 2015), the 420 dynamics were inferred by integrating observations about the magnitude of the displacement across trials. 421 If vestibular cues can in fact support inference of dynamics - as recent findings suggest in eye-head gaze 422 shifts (Sağlam et al., 2014) – a common processing mechanism could be shared across sensory modalities. 423 Overall, this finding highlights the importance of incorporating estimates of the control dynamics in 424 models of self-motion perception and path integration. 425

426 Limitations and future directions

427 Note that restrictions of our motion platform limited the range of velocities that could be tested, allowing

428 only for relatively small velocities (see **Methods**). Consequently, trial durations were long, but the motion 429 platform also restricted total displacement, so we could not test larger target distances. We previously

430 studied visual path integration with larger velocities and our results in the visual and combined conditions

431 are comparable for similar travel times (as trials exceeded durations of 10 seconds, undershooting became 432 more prevalent; Lakshminarasimhan et al., 2018). However, it is unclear how larger velocities (and 433 accelerations) would affect participants' performance (especially under the vestibular condition) and 434 whether the present conclusions are also representative of the regime of velocities not tested.

The design of the Motion Cueing algorithm allowed us to circumvent the issues associated with the 435 physical limitations of the platform to a large degree. This was achieved in part by exploiting the 436 tilt/translation ambiguity and substituting linear translation with tilt (see Methods). However, high 437 frequency accelerations, as those found at movement onset, generated tilts that briefly exceeded the tilt-438 detection threshold of the semicircular canals (Fig. 1 Suppl. 2). Although the duration of suprathreshold 439 stimulation was very small, we cannot exclude the possibility that the perceived tilt affected the 440 interpretation of vestibular inputs. For example, participants may not attribute tilt to linear translation, 441 hence underestimating their displacement. This, however, would lead to overshooting to compensate for 442 the lack of perceived displacement, which is not what we observed in our experiment. Another potential 443 explanation for the poor vestibular performance could be that participants perceive tilt as a conflicting cue 444 with respect to their expected motion or visual cues. In that case, participants would only use the vestibular 445 inputs to a small extent if at all. Manipulating vestibular inputs (e.g. gain, noise manipulations) in future 446 experiments, either alone or in conjunction with visual cues, would offer valuable insights on two fronts: 447 first, to help clarify the efficiency of our Motion Cueing algorithm and its implications on the design of 448 driving simulators in the future, and second, to precisely quantify the contribution of vestibular cues to 449 path integration in natural settings. 450

For the sake of simplicity, we modeled each trial's control dynamics as a single measurement per trial when, in reality, participants must infer the dynamics over the course of a trial using a dynamic process of evidence accumulation. Specifically, participants must measure their self-motion velocity over time and combine a series of measurements to extract information about the underlying dynamics. Although we were able to explain the experimental findings of the influence of control dynamics on steering responses with our model, this approach could be expanded into a more normative framework using hierarchical Bayesian models (Mathys et al., 2011) to infer subjective position estimates by marginalizing over possible control dynamics.

One interesting question is whether providing feedback would eliminate the inference bias of the control 459 dynamics estimation and future studies should explicitly test this hypothesis. Furthermore, it would be 460 interesting to jointly introduce sensory conflict and manipulate sensory reliability to study dynamic 461 multisensory integration such that sensory contributions during navigation can be better disentangled. 462 Although it has been shown that cue combination takes place during path integration (Tcheang et al., 463 2011), previous studies have had contradicting results regarding the manner in which body-based and 464 visual cues are combined (Campos et al., 2010; Chrastil et al., 2019; Koppen et al., 2019; Petrini et al., 465 2016; Horst et al., 2015). Since visual and vestibular signals differ in their sensitivity to different types of 466 motion (Karmali et al., 2014), the outcomes of their integration may depend on the self-motion stimuli 467 employed. Combined with hierarchical models of self-motion inference that considers the control 468 dynamics, it is possible to develop an integrated, multi-level model of navigation, while constraining 469 470 dramatically the hypothesized brain computations and their neurophysiological correlates.

471 Methods

472 EQUIPMENT AND TASK

473 15 participants (9 Male, 6 Female; all adults in the age group 18-32) participated in the experiments. Apart 474 from two participants, all participants were unaware of the purpose of the study. Experiments were first 475 performed in the above two participants before testing others. All experimental procedures were approved 476 by the Institutional Review Board at Baylor College of Medicine and all participants signed an approved 477 consent form.

478 Experimental setup

The participants sat comfortably on a chair mounted on an electric motor allowing unrestricted yaw 479 rotation (Kollmorgen motor DH142M-13-1320, Kollmorgen, Radford, VA), itself mounted on a six-480 degree-of-freedom motion platform (comprised of MOOG 6DOF2000E, Moog Inc., East Aurora, NY). 481 Participants used an analog joystick (M20U9T-N82, CTI electronics, Stratford, CT) with two degrees of 482 freedom and a circular displacement boundary to control their linear and angular speed in a virtual 483 environment based on visual and/or vestibular feedback. The visual stimulus was projected (Canon LV-484 8235 UST Multimedia Projector, Canon USA, Melville, NY) onto a large rectangular screen (width × 485 height: 158×94 cm) positioned in front of the participant (77 cm from the rear of the head). Participants 486 wore crosstalk free ferroelectric active-shutter 3D goggles (RealD CE4s, ColorLink Japan, Ltd., Tokyo, 487 Japan) to view the stimulus. Participants wore headphones generating white noise to mask the auditory 488 motion cues. The participants' head was fixed on the chair using an adjustable CIVCO FirmFit 489 Thermoplastic face mask (CIVCO, Coralville, IA). 490

491 Spike2 software (Power 1401 MkII data acquisition system from Cambridge Electronic Design Ltd., 492 Cambridge, United Kingdom) was used to record joystick and all event markers for offline analysis at a

493 sampling rate of $833\frac{1}{3}$ Hz.

494 Visual stimulus

Visual stimuli were generated and rendered using C++ Open Graphics Library (OpenGL) by continuously 495 repositioning the camera based on joystick inputs to update the visual scene at 60 Hz. The camera was 496 positioned at a height of 70cm above the ground plane, whose textural elements lifetimes were limited 497 $(\sim 250 \text{ ms})$ to avoid serving as landmarks. The ground plane was circular with a radius of 37.5m (near and 498 far clipping planes at 5cm and 3750cm respectively), with the participant positioned at its center at the 499 beginning of each trial. Each texture element was an isosceles triangle (base \times height 5.95 \times 12.95 cm) 500 that was randomly repositioned and reoriented at the end of its lifetime. The floor density was held 501 constant across trials at $\rho = 2.5$ elements/m². The target, a circle of radius 25cm whose luminance was 502 matched to the texture elements, flickered at 5Hz and appeared at a random location between $\theta = \pm 38^{\circ}$ 503 of visual angle at a distance of r = 2.5 - 5.5 m (average distance $\bar{r} = 4$ m) relative to where the 504 participant was stationed at the beginning of the trial. The stereoscopic visual stimulus was rendered in an 505 alternate frame sequencing format and participants wore active-shutter 3D goggles to view the stimulus. 506

507 Behavioral task – Visual, Inertial and Multisensory motion cues

508 Participants were asked to navigate to a remembered target ('firefly') location on a horizontal virtual plane using a joystick, rendered in 3D from a forward-facing vantage point above the plane. Participants pressed 509 510 a button on the joystick to initiate each trial and were tasked with steering to a randomly placed target that was cued briefly at the beginning of the trial. A short tone at every button push indicated the beginning of 511 the trial and the appearance of the target. After one second, the target disappeared, which was a cue for 512 the participant to start steering. During steering, visual and/or vestibular/somatosensory sensory feedback 513 was provided (see below). Participants were instructed to stop at the remembered target location, and then 514 push the button to register their final position and start the next trial. Participants did not receive any 515 feedback about their performance. Prior to the first session, all participants performed about ten practice 516 trials to familiarize themselves with joystick movements and the task structure. 517

The three sensory conditions (visual, vestibular, combined) were randomly interleaved. In the visual condition, participants had to navigate towards the remembered target position given only visual information (optic flow). Visual feedback was stereoscopic, composed of flashing triangles to provide self-motion information but no landmark. In the vestibular condition, after the target disappeared, the entire visual stimulus was shut off too, leaving the participants to navigate in complete darkness using only vestibular/somatosensory cues generated by the motion platform. In the combined condition, participants were provided with both visual and vestibular information during their movement.

525 Independently of the manipulation of the sensory information, the properties of the motion controller also 526 varied from trial to trial. Participants experienced different time constants in each trial, which affected the 527 type and amount of control that was required to complete the task. In trials with short time constants, 528 joystick position mainly controlled velocity, whereas in trials with long time constants, joystick position 529 approximately controlled the acceleration (explained in detail in *Control Dynamics* below).

Each participant performed a total of about 1450 trials (mean \pm standard deviation (SD): 1450 \pm 224), split equally among the three sensory conditions (mean \pm SD – vestibular: 476 \pm 71, visual: 487 \pm 77, combined: 487 \pm 77). We aimed for at least 1200 total trials per participant, and collected extended data from participants whose availability was compatible with the long runtime of our experiment.

534 Joystick control

Participants navigated in the virtual environment using a joystick placed in front of the participant's 535 midline, in a holder mounted on the bottom of the screen. This ensured that the joystick was parallel to 536 the participant's vertical axis, and its horizontal orientation aligned to the forward movement axis. The 537 joystick had two degrees of freedom that controlled linear and angular motion. Joystick displacements 538 were physically bounded to lie within a disk, and digitally bounded to lie within a square. Displacement 539 of the joystick over the anterior-posterior (AP) axis resulted in forward or backward translational motion, 540 whereas displacement in the left-right (LR) axis resulted in rotational motion. The joystick was enabled 541 after the disappearance of the target. To avoid skipping trials and abrupt stops, the button used to initiate 542 trials was activated only when the participant's velocity dropped below 1 cm/s. 543

The joystick controlled both the visual and vestibular stimuli through an algorithm that involved two processes. The first varied the <u>control dynamics (CD)</u>, producing velocities given by a lowpass filtering of the joystick input, mimicking an inertial body under viscous damping. The time constant for the control

547 filter (control timescale) was varied from trial to trial, according to a correlated random process as 548 explained below.

549 The second process was a motion cueing (MC) algorithm applied to the output of the CD process, which

550 defined physical motion that approximated the accelerations an observer would feel under the desired

551 control dynamics, while avoiding the hardwired constraints of the motion platform. This motion cueing

algorithm trades translation for tilt, allowing extended acceleration without hitting the displacement limits(24 cm).

554 These two processes are explained in detail below.

555 Control Dynamics (CD)

Inertia under viscous damping was introduced by applying a lowpass filter on the control input, following an exponential weighted moving average with a time constant that slowly varied across trials. On each trial, the system state evolved according to a first-order Markov process in discrete time, such that the movement velocity at the next time step depended only on the current joystick input and the current velocity. Specifically, the vertical and horizontal joystick positions u_t^v and u_t^ω determined the linear and angular velocities v_t and ω_t as

$$v_{t+1} = \alpha v_t + \beta_v u_t^v$$
 and $\omega_{t+1} = \alpha \omega_t + \beta_\omega u_t^\omega$ (1.1)

562 The time constant τ of the lowpass filter determined the coefficient α (Figure 1 supplement 3A):

$$\alpha$$
 (

$$= e^{-\Delta t/\tau}$$

563

564 Sustained maximal controller inputs of $u_t^v = 1$ or $u_{\omega}^v = 1$ produce velocities that saturate at

$$v_{\max} = \beta_v / (1 - \alpha)$$
 and $\omega_{\max} = \beta_\omega / (1 - \alpha)$ (1.3)

We wanted to set v_{max} and ω_{max} in such a way that would ensure that a target at an average linear or angular displacement x is reachable in an average time T, regardless of τ (we set x = 4 m and T = 8.5 s). This constrains the input gains β_v and β_ω . We derived these desired gains based on a 1D bang-bang control model (*i.e.* purely forward movement, or pure turning) which assumes maximal positive control until time s, followed by maximal negative control until time T (Figure 1 supplement 3A). Although we implemented the leaky integration in discrete time with a frame rate of 60Hz, we derived the input gains using continuous time and translated them to discrete time.

572 The velocity at any time $0 \le t \le T$ during the control is:

$$\frac{v_t}{v_{\max}} = \begin{cases} 1 - e^{-t/\tau} & 0 < t \le s \\ -1 + \left(\frac{v_s}{v_{\max}} + 1\right) e^{-\frac{t-s}{\tau}} & s < t < T \end{cases}$$
(1.4)

573 where v_s is the velocity at the switching time *s* when control switched from positive to negative, given 574 by:

$$v_s = v_{\max} (1 - e^{-s/\tau})$$
 (1.5)

575 By substituting v_s into Eq. (1.4) and using the fact that at time *T*, the controlled velocity should return to 576 0, we obtain an expression that we can use to solve for *s*:

$$v_T = 0 = -1 + \left(\frac{v_{\max}\left(1 - e^{-s/\tau}\right)}{v_{\max}} + 1\right) e^{-\frac{T-s}{\tau}}$$
(1.6)

577 Observe that v_{max} cancels in this equation, so the switching time *s* is independent of v_{max} and therefore 578 also independent of the displacement *x* (see also **Figure 1 supplement 3A**):

$$s = \tau \ln\left(\frac{1 + e^{T/\tau}}{2}\right) \tag{1.7}$$

579 Integrating the velocity profile of Equation 1.4 to obtain the distance travelled by time T, substituting the 580 switch time s (Figure 1 supplement 3A), and simplifying, we obtain:

$$x = x_T = 2 \tau v_{\max} \ln\left(\cosh\frac{T}{2\tau}\right)$$
(1.8)

581 We can then solve for the desired maximum linear speed v_{max} for any time constant τ , average 582 displacement x and trial duration T:

$$v_{\max}(\tau) = \frac{x}{2\tau} \frac{1}{\ln \cosh(T/2\tau)}$$
(1.9)

583 Similarly, the maximum angular velocity was: $\omega_{\max}(\tau) = \frac{\theta}{2\tau} \frac{1}{\ln \cosh(T/t)}$, where θ is the average angle 584 we want our participant to be able to turn within the average time T.

585 These equations can also be re-written in terms of a dimensionless time $z = \tau/T$ (duration of trial in units 586 of the time constant) and average velocities $\bar{v} = x/T$ and $\bar{\omega} = \theta/T$:

$$v_{\max} = \bar{v}_{\frac{1/2z}{\ln\cosh(1/2z)}} \qquad \qquad \omega_{\max} = \bar{\omega}_{\frac{1/2z}{\ln\cosh(1/2z)}} \qquad (1.10)$$

587 where θ is the average angle we want the participants to be able to steer within time T.

588 Setting control gains according to Equation 1.9 allows us to manipulate the control timescale τ , while

⁵⁸⁹ approximately maintaining the average trial duration for each target location (Figure 1 supplement 3B).

590 Converting these maximal velocities into discrete-time control gains using Equations 1.1–1.3 gives us the

591 desired inertial control dynamics.

592 Slow changes in time constant

593 The time constant τ was sampled according to a temporally correlated log-normal distribution. The log of 594 the time constant, $\phi = \log \tau$, followed a bounded random walk across trials according to (Figure 1 595 supplement 3C)

$$\phi_{t+1} = c \phi_t + \eta_t \tag{2}$$

The marginal distribution of ϕ was normal, $N(\mu_{\phi}, \sigma_{\phi}^2)$, with mean $\mu_{\phi} = \frac{1}{2} (\ln \tau_- + \ln \tau_+)$ and standard deviation $\sigma_{\phi} = \frac{1}{4} (\ln \tau_+ - \ln \tau_-)$, which ensured that 95% of the velocity timescales lay between $\tau_$ and τ_+ . The velocity timescales changed across trials with their own timescale τ_{ϕ} , related to the update coefficient by $c = e^{-\Delta t/\tau_{\phi}}$, where we set Δt to be one trial and τ_{ϕ} to be two trials. To produce the desired equilibrium distribution of ϕ we set the scale of the random walk Gaussian noise $\eta \sim N(\mu_{\eta}, \sigma_{\eta}^2)$ with $\mu_{\eta} = \mu_{\phi}(1-c)$ and $\sigma_{\eta}^2 = \sigma_{\phi}^2(1-c^2)$.

602 Motion Cueing algorithm (MC)

Each motion trajectory consisted of a linear displacement in the 2D virtual space combined with a rotation 603 in the horizontal plane. While the motion platform could reproduce the rotational movement using the 604 yaw motor (which was unconstrained in movement range and powerful enough to render any angular 605 acceleration or speed in this study), its ability to reproduce linear movement was limited by the platform's 606 maximum range of 25 cm and maximum velocity of 50 cm/s (in practice, the platform was powerful 607 enough to render any linear acceleration in this study). To circumvent this limitation, we designed a MC 608 algorithm that takes advantage of the gravito-inertial ambiguity (Einstein, 1907) inherent to the vestibular 609 organs (Angelaki & Dickman, 2000; Fernandez et al., 1972; Fernandez & Goldberg, 1976). 610

Specifically, the otolith organs in the inner ear sense both linear acceleration (A) and gravity (G), *i.e.* they 611 sense the gravito-inertial acceleration (GIA): F = G + A. Consequently, a forward acceleration of the 612 head $(a_x, \text{ expressed in g, with } 1g = 9.81 \text{ m/s}^2)$ and a backward pitch (by an angle θ , in radians) will 613 generate a total gravito-inertial acceleration $Fx = \theta + a_x$. The MC took advantage of this ambiguity to 614 replace linear acceleration by tilt. Specifically, it controlled the motion platform to produce a total GIA 615 (Fig. 1 supplement 1, 'Desired Platform GIA') that matched the linear acceleration of the simulated 616 motion in the virtual environment. As long as the rotation that induced this simulated acceleration was 617 slow enough, the motion felt subjectively was a linear acceleration. 618

This control algorithm was based on a trade-off where the high-pass component of the simulated inertial acceleration (**Fig. 1 supplement 1**, 'Desired Platform Linear Acceleration') was produced by translating the platform, whereas the low-pass component was produced by tilting the platform (**Fig. 1 supplement 1**, 'Desired Platform Tilt').

Even though this method is generally sufficient to ensure that platform motion remains within its envelope, 623 it does not guarantee it. Thus, the platform's position, velocity and acceleration commands were fed 624 though a sigmoid function f (Fig. 1 supplement 1, 'Platform Limits'). This function was equal to the 625 identity function (f(x) = x) as long as motion commands were within 75% of the platform's limits, so 626 these motion commands were unaffected. When motion commands exceed this range, the function bends 627 smoothly to saturate at a value set slightly below the limit, thus preventing the platform from reaching its 628 mechanical range (in position, velocity or acceleration) while ensuring a smooth trajectory. Thus, if the 629 desired motion exceeds 75% of the platform's performance envelope, the actual motion of the platform is 630 diminished, such that the total GIA actually experienced by the participant ('Actual Platform GIA') may 631 not match the desired GIA. If left uncorrected, these GIA errors would result in a mismatch between 632 inertial motion and the visual VR stimulus. To prevent these mismatches, we designed a loop that 633

634 estimates GIA error and updates the simulated motion in the visual environment. For instance, if the joystick input commands a large forward acceleration and the platform is unable to reproduce this 635 636 acceleration, then the visual motion is updated to represent a slower acceleration that matches the platform's motion. Altogether, the IC and MC algorithms are applied sequentially as follows: 1) The 637 velocity signal produced by the IC process controls the participant's attempted motion in the virtual 638 environment. 2) The participant acceleration in the VR environment is calculated and inputted to the MC 639 algorithm ('Desired Platform GIA'). 3) The MC cueing computes the platform's motion commands and 640 the actual platform GIA is computed. 4) The difference between the Desired GIA motion actual GIA (GIA 641 error) is computed and used to update the motion in the virtual environment. 5) The updated position is 642 sent to the visual display. 643

A summary of the performance and efficiency of the MC algorithm during the experiment can be seen in **Fig. 1 supplement 2**. For a detailed view of the implementation of the MC algorithm refer to the **Appendix.**

647 QUANTIFICATION AND STATISTICAL ANALYSIS

648 Customized MATLAB code was written to analyze data and to fit models. Depending on the quantity 649 estimated, we report statistical dispersions either using 95% confidence interval, standard deviation, or 650 standard error in the mean. The specific dispersion measure is identified in the portion of the text 651 accompanying the estimates. For error bars in figures, we provide this information in the caption of the 652 corresponding figure. We report and describe the outcome as significant if p < 0.05.

653 Estimation of response gain

In each sensory condition, we first computed the τ -independent gain for each participant; we regressed (without an intercept term) each participant's response positions $(\tilde{r}, \tilde{\theta})$ against target positions (r, θ) separately for the radial (\tilde{r} vs r) and angular ($\tilde{\theta}$ vs θ) coordinates, and the radial and angular response gains (g_r, g_θ) were quantified as the slope of the respective regressions (**Fig 2A**). In addition, we followed the same process to calculate gain terms within three τ groups of equal size (**Fig. 3A**).

659 Correlation between residual error and time constant τ

To evaluate the influence of the time constant on the steering responses, we computed the correlation coefficient between the time constants and the residual errors from the mean response (estimated using the response gain) for distance and angle. Under each sensory condition, the radial residual error (ε_r) for each trial *i* was given by:

$$\varepsilon_{r,i} = \tilde{r}_i - g_r r_i \tag{3.1}$$

where \tilde{r}_i is the radial response, and the mean radial response is given by multiplying the target distance r_i by the radial gain g_r . Similarly, the angular residual error (ε_{θ}) was calculated as:

$$\varepsilon_{\theta,i} = \tilde{\theta}_i - g_\theta \theta_i \tag{3.2}$$

666 Regression model containing τ

⁶⁶⁷ To assess the manner in which the time constant affected the steering responses, we augmented the simple ⁶⁶⁸ linear regression models for response gain estimation mentioned above with τ -dependent terms (Fig. 3

supplement 2; τ and $\tau * r$ for radial response \tilde{r} , τ and $\tau * \theta$ for angular response $\tilde{\theta}$). Subsequently, we calculated the Pearson linear partial correlations between the response positions and each of the three predictors.

672 Estimation of *τ*-dependent gain

To quantify the extent to which the time constant modulates the response gain, we linearly regressed each participant's response positions $(\tilde{r}, \tilde{\theta})$ against target positions (r, θ) and the interaction between target positions and the time constant τ according to:

$$\tilde{r} = b_r r + a_r r \tau$$
 and $\tilde{\theta} = b_\theta \theta + a_\theta \theta \tau$ (4.1)

676 Where b_r , b_θ and a_r , a_θ are the coefficients of the target locations and the interaction terms, respectively. 677 All quantities were first standardized by dividing them with their respective standard deviation, to avoid 678 size effects of the different predictors. This form allows for modulation of the response gain by the time 679 constant, which is clear when the target location is factored out:

$$\tilde{r} = r(b_r + a_r \tau)$$
 and $\tilde{\theta} = \theta(b_\theta + a_\theta \tau)$ (4.2)

680

681 Estimation of simulated no-adaptation response gains

We quantified the extent to which participants failed to adapt to the underlying control dynamics, by 682 generating a simulated null case for no adaptation. First, we selected trials in which the time constant was 683 close to the mean of the sampling distribution $(\pm 0.2s)$. Then, we integrated the steering input of those trials 684 with time constants from other trials (see equations 1.1, 1.2). This generated a set of trajectories for which 685 the steering corresponded to a different time constant, providing us with a null case of no adaptation to 686 the underlying dynamics. We then stratified the simulated trajectories into equal-sized groups based on 687 the time constants (same as in Fig. 3A) and computed the corresponding radial and angular response gains. 688 Note that the response gains were computed according to the target locations of the initial set of trials. 689

690 Rationale behind modeling approach

We tested the hypothesis that the τ -dependent errors in steering responses arise from participants 691 misestimating control dynamics on individual trials. Specifically, if participants' estimate of the time 692 constant τ differs from the actual value, then their *believed* trajectories (computed using the estimated τ) 693 would differ accordingly from the actual trajectories along which they travelled. believed stopping 694 locations should land on or near the target. However, various unmeasurable fluctuations in that belief 695 across trials should lead to variability clustered around the target location Because participants try to stop 696 on their believed target location, the believed stopping locations, subject to unmeasurable fluctuations of 697 698 the belief across trials, should be distributed evenly around the participant's mean response (mean belief), after adjusting for the average response gain. This is so because, if the distribution of believed responses 699 depended on the time constant, then that would imply that participants willingly misestimated the control 700 dynamics. Mathematically, the subjective residual errors (deviation of the believed stopping location from 701 the mean response for a given target; see Methods: Correlation between residual error and time constant 702 703 τ) should be distributed evenly around zero and be uncorrelated with the time constant τ . Therefore, a good

model of the participants' beliefs should predict that subjective residual errors are statistically independentof the time constant.

706 **Bayesian observer model for** τ **estimation**

707 To account for the effect of the time constant τ on steering performance, we considered a two-step observer model that uses a measurement m of the real time constant τ and a prior distribution over hypothesized time constants in logarithmic scale to compute an estimate $\hat{\tau}$ on each trial (first step), and then integrates 709 the actual joystick input using that estimate to reconstruct the participant's believed trajectory (second 710 711 step). We formulated our model in the logarithmic space of $\varphi = \log \tau$, therefore the prior distribution over the hypothesized time constants $p(\varphi)$ was assumed to be normal in log-space with mean, μ_{prior} and standard deviation, σ_{prior} . The measurement distribution $p(m|\varphi)$ was also assumed to be normal in log-713 space with mean φ , and standard deviation σ_{measure} . Note that whereas the prior $p(\varphi)$ remains fixed 714 across trials of a particular sensory modality, the mean of measurement distribution is governed by φ and 715 716 thus varies across trials. For each sensory modality, we fit two parameters, $\Theta \ni \{\mu_{\text{prior}}, \lambda\}$, where λ was taken to be the ratio of σ_{prior} over σ_{measure} , (i.e. their relative weight). 717

718 Model fitting

719 When inferring the participant's beliefs about the control dynamics, we computed the posterior 720 distribution on trial *i* as $p(\varphi|m_i) \propto p(\varphi)p(m_i|\varphi)$ (Fig. 5A, left) and then selected the median over φ 721 (equal to the maximum a posteriori estimate), and back-transformed it from log-space to obtain an estimate 722 of the time constant $\hat{\tau}_i$ for that trial:

$$\hat{\tau}_i = \exp\left\{ \underset{\varphi}{\operatorname{argmax}} p(\varphi | m_i) \right\}$$
(5)

723 Subsequently, $\hat{\tau}$ is used to integrate the actual joystick input and produce the participant's believed 724 trajectory, according to (Equation 1.1-1.10) in the *Control Dynamics* (CD) section.

The Bayesian model had 2 free parameters $\Theta \ni \{\mu_{\text{prior}}, \lambda\}$. We fit the model by assuming that participants stop as close to the target as possible given their understanding of the task. Specifically, we minimized the mean squared error (MSE) between the measured mean stopping position (computed using the response gains g_r and g_{θ} from **Equation 3**) and our model of the participant's believed stopping location $\hat{\mathbf{x}}_i$ given the inferred dynamics $\hat{\tau}_i$. For each sensory condition:

$$\Theta^* = \underset{\Theta}{\operatorname{argmin}} \quad \frac{1}{n} \sum_{i=1}^n \{ \hat{\mathbf{x}}_i(\hat{\tau}_i, \mathbf{u}_i) - \mathbf{G} \, \mathbf{x}_i^{\operatorname{tar}} \}^2 \tag{6}$$

where, for each trial i, $\hat{\mathbf{x}}_i$ is the believed participant's position, $\hat{\tau}_i$ is the estimated time constant, \mathbf{u}_i is the time series of the joystick control input, $\mathbf{x}_i^{\text{tar}}$ is the actual target position, **G** is the response gain matrix determined from g_r and g_{θ} , and n is the total number of trials.

733 Model validation

734 To evaluate the performance of our model, we examined the correlations between the *subjective* residual error and τ that are given by the model. The subjective residual error is defined as the difference between 735 the believed (subjective) stopping location that a model produces and the mean response of the actual 736 trajectories, adjusted for the response gain. The subjective residual errors are calculated for the radial and 737 angular components of the response separately, according to Equation 3 (where actual responses $\tilde{r}, \tilde{\theta}$ are 738 substituted by believed responses $\hat{\tilde{r}}, \hat{\tilde{\theta}}$, respectively). Ideally, these correlations should not exist for the 739 model predictions (explained in text; Fig. 5B). We determined the statistical significance of the model-740 implied correlations by adjusting for multiple comparisons (required level of statistical significance: 741 p=0.0085). To assess the performance of the Bayesian model, we compared the correlations between 742 believed and actual stopping location with the time constant (Fig. 5B; Wilcoxon signed-rank test). 743

744 Dynamic prior model

Since the time constant changes randomly across trials, we tested whether the history of time constants influenced the estimate $\hat{\tau}$. If true, the Bayesian model would imply a prior distribution over $\varphi = \log \tau$ that is dynamically changing according to the recent history of time constants, rather than being fixed. To explore this possibility, we repeated the two-step model outlined above, with the difference that the mean of the prior distribution is updated at every trial *i* by a weighted average of the mean prior in the previous trial and the current measurement over φ :

$$\mu_{\text{prior},i} = (1-k)\,\mu_{\text{prior},i-1} + k\,\varphi_i \text{ where } k = \frac{\lambda^2}{\lambda^2 + 1}$$
(7)

and where λ is the ratio of prior standard deviation over likelihood standard deviation. As *k* indicates, the relative weighting between prior and measurement on each trial depends solely on their relative widths. Finally, the initial prior was taken to be the time constant on the first trial. Thus, the only free parameter we fit was λ .

755 Sensory-independent model

As another alternative to the Bayesian model with a static prior, we also constructed a model where participants ignored changes in the time constants and navigated according to a fixed estimate $\hat{\tau}$ across all trials in each sensory condition. This model had only one free parameter: the time constant estimate $\hat{\tau}$, which was integrated with the actual joystick input of each trial to reconstruct the believed trajectory of the participant. We fit $\hat{\tau}$ for each sensory condition by minimizing the MSE between the believed stopping location and the mean response (according to **Equation 6**).

762 Model comparison

763 To compare the static prior Bayesian model against the dynamic prior Bayesian and the sensory-764 independent models, we compared the correlations between believed stopping locations and time 765 constants that each model produces (**Fig. 7**; paired Student's t-test).

766 Sensory feedback control model

We tested a sensory feedback control model, in which the controller uses bang-bang control and switches from forward to backward input at a constant and predetermined distance from the target position (corrected for the bias, *i.e.* mean response). Specifically, we preserved the actual angular and only fitted the linear control input for each trial. Thus, as switch distance, we refer to a Euclidian distance from the bias-corrected target position. We fit the mean and standard deviation of the switch distance for each participant in each condition separately, by minimizing the distance of the actual from the model-predicted stopping locations. To evaluate how well this model describes our data, we compared the correlations and regression slopes between the time constant and residual errors from the stopping locations predicted by the model with those from our actual data (**Fig. 7 supplement 2**).

776

777 DATA AND SOFTWARE AVAILABILITY

778 MATLAB code implementing all quantitative analyses in this study is available online 779 (<u>https://github.com/ AkisStavropoulos/matlab_code</u>). Datasets generated by this study are available online 780 (<u>https://gin.g-node.org/akis stavropoulos/humans control dynamics sensory modality steering</u>).

781 Acknowledgements

We thank Jing Lin and Jian Chen for their technical support, and Baptiste Caziot, Panos Alefantis, Babis
Stavropoulos and Evangelia Pappou for their useful insights. This work was supported by the Simons
Collaboration on the Global Brain, grant no. 324143, and NIH DC007620. G.C.D. was supported
by NIH EY016178.

786 Competing interests

787 The authors declare no competing interests.



Figure 1 supplement 1: (A) Flow diagram of Motion Cueing algorithm (MC). The participant pilots themselves in a simulated environment using a joystick. The motion cueing algorithm aims at controlling a platform such that the sum of inertial and gravitational acceleration experienced when sitting on the platform (*desired platform GIA*, blue; the curve illustrates an example profile consisting of a single rectangular waveform) matches the linear acceleration experienced in the simulated virtual environment. "Desired" refers to the fact that the motion platform may not be able to match this acceleration exactly. The desired GIA is fed through a step impulse function to compute the *desired linear acceleration* of the platform. The difference between the desired linear acceleration and GIA is used to compute the *desired platform tilt*. The desired platform motion (linear and tilt motion) are passed through a controller that restricts its motion to the actuator's limits (in term of linear and angular acceleration, velocity, and position). The two actuator output commands are sent to the platform and are also used to compute the actuator GIA which is actually rendered by the platform. To ensure that the inertial motion produced by the platform matches the motion in the simulated environment, the actuator GIA is compared to the desired linear acceleration to compute an actuator GIA error feedback signal, which updates the simulated motion. (B) Acceleration profile of an actual trial. The first panel shows the desired GIA of the participant for that trial. The second and third panels show the desired linear acceleration (red) and desired tilt acceleration (green), respectively. The fourth panel shows the final GIA achieved (blue) and the GIA error (magenta). (C) Correspondence between visual acceleration and platform GIA (blue), measured independently from the motion cueing algorithm using an inertial measurement unit mounted next to the participant's head. There is an almost perfect match between the two. The gray histogram indicates the range of acceleration experienced by the participant.





Figure 1 supplement 2: (A) Net Gravito-Inertial Acceleration (GIA; thick lines) and net GIA error (thin lines) aligned to start and end of trial, for the vestibular and combined conditions across participants (average across trials over all time constants). The dashed line represents a conservative choice of the vestibular motion detection threshold according to the relevant literature (8 cm/s²; Kingma, 2005; MacNeilage et al., 2010; Zupan & Merfeld, 2008). Gray region represents the target presentation period. Shaded regions denote ± 1 SEM. (B) Net tilt velocity aligned to start and end of trial, for the vestibular and combined conditions across participants. Dashed line represents the estimated tilt/translation discrimination threshold of 1 deg/s: although tilt/translation discrimination thresholds have not been explicitly studied, we can use the rotation sensation thresholds of the semicircular canals to estimate what that threshold would be. Since it is the rotation velocity that tells a participant that they are tilting and not translating, we propose that the tilt/translation discrimination threshold is at least the same as the rotation sensation threshold (if not larger; Lim et al., 2017; MacNeilage et al., 2010). Shaded regions represent ± 1 SEM across participants. Inset shows the probability distribution of displacements during the supra-threshold tilt period after trial onset (~ 0.6 s). Although the tilt can be perceived by the participants during trial onset, the displacement during that period does not exceed 10cm and could potentially not contribute significantly to steering errors, for three reasons: a) the displacement during that period is negligible, b) tilt velocity is kept below the perceptual threshold for the remainder of the trajectory, c) GIA is always above the motion detection threshold of the vestibular system. However, since the initial tilt could be perceived (as it briefly exceeded the canal detection threshold), this might alter the perceived orientation of the participants. In turn, this could influence the extent to which vestibular cues would be used as input to the path integration system (see Discussion "Limitations and future directions" for further discussion).

Thus, perceived tilt might be used as an indicator of trial onset, but it cannot contribute to path integration for 3 reasons: a) the displacement during that period is negligible, b) tilt velocity is kept below the perceptual threshold for the remainder of the trajectory, c) GIA is always above the motion detection threshold of the vestibular system.



Figure 1 supplement 3: (A) Example dynamics for bang-bang control. Position, velocity, and controls are shown. Control switches at time *s* and ends at time *T*. (B) Maximal velocity (blue) needed for bang-bang control to produce a desired average velocity $\bar{u} = x/T$, as a function of the fraction of trial duration given by the time constant, τ/T . When the time constant is a small fraction of the trial (velocity control), the max velocity equals the average velocity (orange line). When the time constant is much longer than a trial (acceleration control), the maximum velocity grows as $4\tau/T$ (green), although this speed is never approached since braking begins before the velocity approaches equilibrium. (C) Example dynamics for control behavior. *Left*: log-normal distribution of control time constants τ (see also Fig. 3 supplement 1A). *Right*: example random walk in log τ space.



Figure 2 supplement 1: (A) Random subset of trajectories of an example participant under each sensory condition. The corresponding radial and angular response gains are indicated for each condition (green: vestibular, cyan: visual, purple: combined). *Gray region* represents the target range. **(B)** Sex differences in participants' performance: radial and angular gains (see **Fig. 2D**) grouped based on sex (F: female, M: male; see legend; *p*-values of differences in response gains between male and female participants: Radial gain – vestibular: p = 0.17, visual: p = 0.09, combined: p = 0.09; Angular gain – vestibular: p = 0.58, visual: p = 0.38, combined: p = 0.21; two-sample *t*-test). **(C)** Sex differences in participants' performance: correlation coefficients between the time constant and the residual errors (radial and angular components; see **Fig.3C**) grouped based on sex. Specifically, the x and y axes represent the correlation values between the time constant and the radial and angular residual errors, respectively (*p*-values of differences in correlation coefficients between male and female participants: Radial – vestibular: p = 0.5, visual: p = 0.66, combined: p = 0.71; Angular – vestibular: p = 0.51, visual: p = 0.97, combined: p = 0.82; two-sample *t*-test).

792



Figure 2 supplement 2: (A-G) Linear (left) and angular (right) joystick input over time, for a subset of participants in all conditions (see *legend*; bottom right). The joystick control had shorter duration in the vestibular condition, reflecting our findings of the smaller response gains. Shaded regions represent ± 1 standard deviation across trials.



Figure 3 supplement 1: (A) Sampling distributions of the time constant for all three sensory conditions across participants. The sampling distribution both across participants and across conditions are almost identical. *Transparent* lines and *thick* lines represent the individual sampling distributions of participants and their mean, respectively. (B-J) Effect of the time constant on radial (left) and angular (right) residual error, for a large subset of participants. *Solid lines* represent linear regression fits (see Table 3 for individual regression coefficient values). *Dashed lines* denote zero residual error (*i.e.* stopping location matches mean response).



Figure 3 supplement 2: (A) Partial correlation coefficients for prediction of stopping distance \tilde{r} (relative to starting position) from initial target distance (r), τ , and the interaction of the two $(r\tau)$, for all participants across sensory conditions. Values at each bar group represent the average coefficient value across participants ±1 standard deviation. The contribution of the τ -only term was considered insignificant across all conditions. The simplified version of this model would be: $\tilde{r} = r(\alpha + \gamma \tau)$, which implies that the radial gain is τ -dependent. (B) Partial correlation coefficients for prediction of stopping angle $\tilde{\theta}$ (relative to starting position) from initial target angle (θ) , τ , and the interaction of the two $(\theta\tau)$, for all participants across sensory conditions. Values at each bar group represent the average coefficient value across participants ±1 standard deviation. In agreement with the findings for the response distance, the contribution of the τ -only term was considered insignificant across all conditions. The simplified version of this model would be: $\tilde{\theta} = \theta(\alpha + \gamma \tau)$, which implies that the angular gain is also τ -dependent.





Figure 5 supplement 1: (A) Correlation coefficients between the time constant and travel duration (**left**) or average travel velocity (**right**) across trials for all participants. *Colors of circles* indicate the sensory condition (*green*: vestibular, *cyan*: visual, *purple*: combined). *Open and filled circles* denote statistical significance according to the legend. (**B**) Dependence of travel duration (*top right*) and average velocity (*bottom right*) on the time constant for perfect or no estimation/adaptation to the dynamics (*left*), for a simulated bang-bang controller. Correlation coefficients and statistical significance are indicated in the legends of the corresponding panels. Solid lines represent linear regression fits. (**C**) **Left:** Uncertainty (variance) of instantaneous self-motion velocity estimation. Illustration of a linear (*blue*) and a quadratic (*orange*) model of velocity estimation uncertainty as a function of the instantaneous velocity magnitude. We wanted to test whether the effect of the time constant on performance could be attributed to differences in the accumulated uncertainty for the linear and quadratic models. We found that the accumulated uncertainty is positively correlated with the time constant for both models (adding an intercept term to the models did not qualitatively change the results). This means that higher time constants yield larger uncertainty and, therefore, participants should undershoot more. However, this is the opposite of the observed effect of the time constant on the responses. Error bars denote ± 1 SEM.





Figure 5 supplement 2: Changes in travel distance for a given control input under different control dynamics. Whether in the domain of velocity (*top*) or acceleration control (*bottom*), a control input that is appropriate to reach a certain target distance (*horizontal black dashed line*) under only a certain time constant (*red vertical line*) will produce erroneous displacements under any other time constant (*blue line*). For smaller time constants, the intended distance will be undershot, whereas larger time constants will lead to overshooting. In other words, assuming that the *red vertical line* denotes the believed dynamics of a controller, a larger actual time constant (underestimation) will lead to overshooting (relative to the intended displacement; *horizontal black dashed line*). Inversely, overestimation of the time constant would lead to undershooting. Note that, for acceleration control we chose a bangbang controller such that we can demonstrate that this holds true whether there is braking at the end of the trial or not.



Figure 7 supplement 1: Correlation coefficients in the vestibular condition between the actual time constant and the subjective radial (left) and angular (right) residual errors, if participants carried over their τ estimate from the previous trial. With sensory conditions interleaved and a common random walk of τ (see Methods, Figure 1 supplement 3C), we searched for a trial-type history effect in the vestibular condition, due to participants' poor τ estimation performance. Specifically, we asked whether participants in the vestibular condition would leverage from the correlation structure between recent time constants by carrying over their estimates from the previous visual or combined trials. We first compared the correlations from the actual data (open bars; same as Fig. 3C, 5B) with those obtained when using the actual (middle bar couple) or estimated (right bar couple; estimates from the static prior Bayesian model) time constant from the previous visual (cvan bars) or combined (purple bars) trial to generate believed trajectories. Although correlations were significantly smaller for the carry-over models relative to the actual data (p < 0.01) they nevertheless remained significant ($p < 10^{-5}$), thus, failing to explain away the effect (compare with grey bars: correlations implied by estimation in the current vestibular trial with the static prior Bayesian model). The carry-over strategy does not seem likely since it fails to explain away a large part of the correlation between the radial component of the subjective residual errors and the time constant (compare rightmost *cyan/purple bars* with *grey bars*; p-values of paired t-test between radial correlation coefficients – current vestibular vs previous visual trial estimation: p=0.006, current vestibular vs previous combined trial estimation: p=0.02; p-values of paired t-test between angular correlation coefficients - current vestibular vs previous visual trial estimation: p=0.008, current vestibular vs previous combined trial estimation: p=0.71). Error bars denote ± 1 SEM across participants.





Figure 7 supplement 2: (A) We tested a sensory feedback control model, in which the controller uses bang-bang control and switches from forward to backward input at a constant and predetermined distance from the target position (corrected for the bias). We fit the mean and standard deviation of the switch distance for each participant in each condition separately, by minimizing the distance of the actual from the model-predicted stopping locations (see **Methods**). The correlations (*left*) and the regression slopes (*right*) between the model-predicted residual errors and the time constant were significantly higher than those found in our data (p-values of difference in correlations between true data and model obtained by paired t-test – vestibular: $p = 10^{-5}$, visual: $p = 10^{-6}$, combined: $p = 10^{-7}$; p-values of difference regression slopes between true data and model obtained by paired t-test – vestibular: $p = 10^{-6}$, combined: $p = 10^{-7}$; visual: $p = 10^{-8}$, combined: $p = 10^{-9}$). *Error bars* represent ±1 SEM across participants. (**B**) Probability distribution of bang-bang switch distance from target position (corrected for the bias). According to the sensory feedback control model, the probability distribution of switch distance should be very narrow since participants switch at a constant perceived distance from the target. If participants implemented this type of control (*black lines*), we would expect to see such a narrow distribution in the actual data. In all conditions, however, the switch distance distribution of the true data (*colored lines*) is wider and resembles what we expect to see if participants implemented optimal (ideal) bang-bang control (*gray lines*). *Shaded regions* represent ±1 SEM across participants.

	Radial Bias Table		
	vestibular	visual	combined
τ: [0.34 - 1.53]	0.649 ± 0.056	0.818 ± 0.057	0.786 ± 0.055
τ: [1.53 - 2.16]	0.733 ± 0.063	0.871 ± 0.059	0.836 ± 0.056
τ: [2.16 - 8.89]	0.902 ± 0.077	0.944 ± 0.061	0.917 ± 0.058

Angular Bias Table

	vestibular	visual	combined
τ: [0.34 - 1.53]	0.731 ± 0.053	0.919 ± 0.036	0.902 ± 0.032
τ: [1.53 - 2.16]	0.770 ± 0.060	0.984 ± 0.038	0.944 ± 0.029
τ: [2.16 - 8.89]	0.878 ± 0.061	1.024 ± 0.040	1.012 ± 0.033

Table 1: Average radial (top) and angular (bottom) behavioral response gains across participants, for groups of time constant τ magnitudes (mean ± SEM).

800

Radial Correlations			
	vestibular	visual	combined
Subject 1	r = 0.585, p = 4.2·10 ⁻⁴⁵	r = 0.502, p = 1.2·10 ⁻³⁷	r = 0.617, p = 1.0·10 ⁻⁵⁹
Subject 2	r = 0.622, p = 5.5·10 ⁻⁴³	r = 0.338, p = 7.4·10 ⁻¹²	r = 0.377, p = 2.9·10 ⁻¹⁴
Subject 3	r = 0.433, p = 3.5·10 ⁻²⁵	r = 0.280, p = 2.7·10 ⁻¹¹	r = 0.374, p = 8.7·10 ⁻²⁰
Subject 4	r = 0.492, p = 9.1·10 ⁻³¹	r = 0.494, p = 3.1·10 ⁻³¹	r = 0.350, p = 2.1·10 ⁻¹⁵
Subject 5	r = 0.411, p = 4.4·10 ⁻¹⁷	r = 0.314, p = 3.4·10 ⁻¹⁰	r = 0.360, p = 3.7·10 ⁻¹³
Subject 6	r = 0.601, p = 2.0·10 ⁻⁵⁸	r = 0.233, p = 1.2·10 ⁻⁰⁸	r = 0.233, p = 1.2·10 ⁻⁰⁸
Subject 7	r = 0.606, p = 1.6·10 ⁻⁴⁴	r = 0.522, p = 1.5·10 ⁻³¹	r = 0.474, p = 1.1·10 ⁻²⁵
Subject 8	r = 0.477, p = 9.6·10 ⁻³⁴	r = 0.255, p = 5.7·10 ⁻¹⁰	r = 0.294, p = 4.6·10 ⁻¹³
Subject 9	r = 0.478, p = 1.0·10 ⁻²²	r = 0.517, p = 7.9·10 ⁻²⁷	r = 0.523, p = 6.3·10 ⁻²⁸
Subject 10	r = 0.573, p = 7.2·10 ⁻³⁹	r = 0.497, p = 4.7·10 ⁻²⁸	r = 0.576, p = 3.5·10 ⁻³⁹
Subject 11	r = 0.375, p = 5.9·10 ⁻¹⁶	r = 0.224, p = 2.1·10 ⁻⁰⁶	r = 0.144, p = 0.002
Subject 12	r = 0.522, p = 2.1·10 ⁻³⁹	r = 0.341, p = 1.3·10 ⁻¹⁶	r = 0.319, p = 1.1·10 ⁻¹⁴
Subject 13	r = 0.512, p = 1.1·10 ⁻³⁸	r = 0.385, p = 1.4·10 ⁻²¹	r = 0.401, p = 4.7·10 ⁻²³
Subject 14	r = 0.461, p = 8.3·10 ⁻³⁰	r = 0.241, p = 1.3·10 ⁻⁰⁸	r = 0.276, p = 7.0·10 ⁻¹¹
Subject 15	r = 0.703, p = 1.7·10 ⁻⁶¹	r = 0.214, p = 1.1·10 ⁻⁰⁵	r = 0.213, p = 1.3·10 ⁻⁰⁵

Angular Correlations

	vestibular	visual	combined
Subject 1	r = 0.254, p = 1.9·10 ⁻⁰⁸	r = 0.302, p = 1.8·10 ^{−13}	r = 0.437, p = 2.3·10 ⁻²⁷
Subject 2	r = 0.156, p = 0.002	r = 0.287, p = 8.6·10 ⁻⁰⁹	r = 0.270, p = 9.2·10 ⁻⁰⁸
Subject 3	r = 0.301, p = 2.2·10 ⁻¹²	r = 0.274, p = 7.3·10 ⁻¹¹	r = 0.351, p = 1.7·10 ⁻¹⁷
Subject 4	r = 0.315, p = 1.3·10 ⁻¹²	r = 0.299, p = 1.7·10 ⁻¹¹	r = 0.343, p = 8.9·10 ⁻¹⁵
Subject 5	r = 0.153, p = 0.003	r = 0.291, p = 6.7·10 ⁻⁰⁹	r = 0.387, p = 3.8·10 ⁻¹⁵
Subject 6	r = 0.292, p = 5.9·10 ⁻¹³	r = 0.121, p = 0.003	r = 0.224, p = 4.8·10 ⁻⁰⁸
Subject 7	r = 0.098, p = 0.042	r = 0.356, p = 2.4·10 ⁻¹⁴	r = 0.275, p = 6.0·10 ⁻⁰⁹
Subject 8	r = 0.346, p = 2.0·10 ⁻¹⁷	r = -0.004, p = 0.920	r = 0.005, p = 0.902
Subject 9	r = 0.093, p = 0.071	r = 0.349, p = 4.1·10 ⁻¹²	r = 0.348, p = 3.1·10 ⁻¹²
Subject 10	r = 0.294, p = 4.7·10 ⁻¹⁰	r = 0.336, p = 9.6·10 ⁻¹³	r = 0.235, p = 9.0·10 ⁻⁰⁷
Subject 11	r = 0.064, p = 0.183	r = -0.032, p = 0.507	r = 0.027, p = 0.575
Subject 12	r = 0.271, p = 1.2·10 ⁻¹⁰	r = 0.278, p = 2.7·10 ⁻¹¹	r = 0.333, p = 5.6·10 ⁻¹⁶
Subject 13	r = 0.238, p = 1.2·10 ⁻⁰⁸	r = 0.312, p = 2.5·10 ⁻¹⁴	r = 0.255, p = 1.0·10 ⁻⁰⁹
Subject 14	r = 0.215, p = 4.3·10 ⁻⁰⁷	r = 0.138, p = 0.001	r = 0.217, p = 3.7·10 ⁻⁰⁷
Subject 15	r = 0.328, p = 1.2·10 ⁻¹¹	r = 0.134, p = 0.006	r = 0.137, p = 0.005

Table 2: Pearson's correlation coefficient (r) and corresponding p-value (p) for radial (top) and angular (bottom) correlation between residual error and the time constant τ across participants. Mean Pearson's $r \pm$ SEM: Radial component – vestibular: 0.52±0.02, visual: 0.36±0.03, combined: 0.37±0.03; Angular component – vestibular: 0.23±0.02, visual: 0.26±0.03.

802

	Radial Reg	ression Coefficient	ts (m/s)
	vestibular	visual	combined
Subject 1	α = 0.775	α = 0.247	α = 0.337
Subject 2	α = 0.776	α = 0.464	α = 0.470
Subject 3	α = 0.255	α = 0.138	α = 0.157
Subject 4	α = 0.406	α = 0.138	α = 0.269
Subject 5	α = 1.009	α = 0.559	α = 0.487
Subject 6	α = 0.829	α = 0.151	α = 0.149
Subject 7	α = 0.512	α = 0.351	α = 0.330
Subject 8	α = 0.582	α = 0.245	α = 0.222
Subject 9	α = 0.321	α = 0.330	α = 0.311
Subject 10	α = 0.943	α = 0.365	α = 0.445
Subject 11	α = 0.522	α = 0.322	α = 0.177
Subject 12	α = 0.484	α = 0.166	α = 0.210
Subject 13	α = 0.570	α = 0.324	α = 0.327
Subject 14	α = 0.507	α = 0.253	α = 0.321
Subject 15	α = 0.799	α = 0.091	α = 0.102

Angular Regression Coefficients (deg/s)

	vestibular	visual	combined
Subject 1	β = 1.664	β = 1.045	β = 1.553
Subject 2	β = 1.645	β = 2.022	β = 1.632
Subject 3	β = 1.317	β = 0.552	β = 1.232
Subject 4	β = 2.165	β = 0.919	β = 1.155
Subject 5	β = 2.349	β = 3.201	β = 3.045
Subject 6	β = 2.620	β = 0.563	β = 0.870
Subject 7	β = 1.434	β = 1.101	β = 0.843
Subject 8	β = 4.185	β = -0.039	β = 0.040
Subject 9	β = 1.254	β = 1.562	β = 1.394
Subject 10	β = 2.937	β = 1.971	β = 1.152
Subject 11	β = 1.849	β = -0.193	β = 0.194
Subject 12	β = 1.382	β = 0.836	β = 0.954
Subject 13	β = 1.619	β = 1.233	β = 1.165
Subject 14	β = 2.141	β = 0.585	β = 0.790
Subject 15	β = 2.214	β = 0.256	β = 0.264

Table 3: Linear regression slope coefficients for radial (α , top) and angular (β , bottom) components of residual error against the time constant τ across participants. Mean regression slope \pm SEM: Radial (m/s) – vestibular: 0.62 \pm 0.06, visual: 0.28 \pm 0.03, combined: 0.29 \pm 0.03; Angular (deg/s) – vestibular: 2.05 \pm 0.2, visual: 1.04 \pm 0.23, combined: 1.09 \pm 0.19.

Radial Partial correlation coefficients ± standard deviation				
		vestibular	visual	combined
Predictors	radial distance (r)	0.20 ± 0.05	0.48 ± 0.13	0.45 ± 0.10
	time constant (τ)	-0.06 ± 0.07	0.01 ± 0.06	-0.03 ± 0.06
	interaction term (r×τ)	0.20 ± 0.09	0.07 ± 0.06	0.12 ± 0.09

Angular Partial correlation coefficients ± standard deviation

		vestibular	visual	combined
Predictors	angular distance (θ)	0.57 ± 0.13	0.90 ± 0.08	0.90 ± 0.06
	time constant (τ)	-0.06 ± 0.08	-0.01 ± 0.06	-0.07 ± 0.06
	interaction term ($\theta \times \tau$)	0.27 ± 0.11	0.28 ± 0.15	0.33 ± 0.14

Table 4: Partial correlation coefficients (mean \pm standard deviation) for prediction of the radial (\tilde{r} , top) and angular ($\tilde{\theta}$, bottom) components of the final stopping location (relative to starting position) from initial target distance (r) and angle (θ), the time constant τ , and the interaction of the two ($r \times \tau$ or $r \times \theta$), respectively.

803

804 Appendix: Implementation of MC algorithm

805 **STEP 1**:

806 In the first step, the participant's velocity is being transformed into the VR (screen) coordinates. This 807 transformation is necessary to deduce centrifugal components from the participants' trajectory, and 808 include them in the motor commands:

$$v_{t+1}^{\text{VR},x} = v_{t+1}^{\text{JS}} \cdot \cos(\varphi_t^{\text{VR}})$$

$$v_{t+1}^{\mathrm{VR}, y} = v_{t+1}^{\mathrm{JS}} \cdot \sin(\varphi_t^{\mathrm{VR}})$$

811

810

812 Where $v^{VR,x}$ and $v^{VR,y}$ is the linear velocity of the participant in VR coordinates, ω^{VR} is the angular 813 velocity of the VR system, and φ^{VR} is the direction of the platform in space.

 $\omega_{t+1}^{\text{VR}} = \omega_{t+1}^{\text{JS}}$

814 STEP 2:

As mentioned before, the arena diameter is finite, and it is necessary to keep track of the participant's position in the arena, to avoid "crashing" on the invisible walls. In this step, the participant's velocity is slowed down when the participant approaches the boundaries of the arena, to account for a "smooth crash".

818 **STEP 3**:

Here, the current acceleration is calculated in the VR coordinates $(a^{VR,x}, a^{VR,y})$. This is also where the GIA error feedback loop (see STEP 10) updates the VR acceleration.

821
$$\alpha_{t+1}^{\text{VR},x} = \frac{\nu_{t+1}^{\text{VR},x} - \nu_t^{\text{VR},x} + \frac{dt}{\tau_{\text{MC}}} \cdot (\nu_t^{\text{VR},x} - \hat{\nu}_t^{\text{VR},x})}{dt}$$

822
$$\alpha_{t+1}^{\text{VR},y} = \frac{v_{t+1}^{\text{VR},y} - v_t^{\text{VR},y} + \frac{dt}{\tau_{\text{MC}}} \cdot (v_t^{\text{VR},y} - \hat{v}_t^{\text{VR},y})}{dt}$$

823

824 Where \hat{v}_t is the updated velocity from the previous timestep (τ_{MC} explained in STEP 10). After the 825 acceleration is obtained, it is being transformed back to the participant's coordinates ($a^{sub,x}, a^{sub,y}$):

826
$$\alpha_{t+1}^{\operatorname{sub},x} = \alpha_{t+1}^{\operatorname{VR},x} \cdot \cos(\varphi_t^{\operatorname{VR}}) + \alpha_{t+1}^{\operatorname{VR},y} \cdot \sin(\varphi_t^{\operatorname{VR}})$$

827
$$\alpha_{t+1}^{\operatorname{sub},y} = -\alpha_{t+1}^{\operatorname{VR},x} \cdot \sin(\varphi_t^{\operatorname{VR}}) + \alpha_{t+1}^{\operatorname{VR},y} \cdot \cos(\varphi_t^{\operatorname{VR}})$$

828 <u>STEP 4:</u>

Now, the acceleration a^{sub} in participant's coordinates, is being transformed into Platform coordinates to take into account the orientation of the participant onto the motion platform (φ_t^{moog}), which is controlled by the yaw motor. For instance, if the participant faces towards the left of the platform and accelerates forward in egocentric coordinates, then the platform should move to the left:

833
$$\alpha_{t+1}^{\text{desired},x} = \alpha_{t+1}^{\text{sub},x} \cdot \cos(\varphi_t^{\text{moog}}) - \alpha_{t+1}^{\text{sub},y} \cdot \sin(\varphi_t^{\text{moog}})$$

834
$$\alpha_{t+1}^{\text{desired},y} = \alpha_{t+1}^{\text{sub},x} \cdot \sin(\varphi_t^{\text{moog}}) + \alpha_{t+1}^{\text{sub},y} \cdot \cos(\varphi_t^{\text{moog}})$$

835 Where $\alpha_{t+1}^{\text{desired},x}$ is the desired platform acceleration.

836 STEP 5:

837 This is the Motion-Cueing (MC) step. Here, the amount of tilt and translation that will be commanded is computed, based on the tilt-translation trade-off we set. First, the platform's desired acceleration is 838 computed by applying a step response function f(t) to the acceleration input: 839

840
$$a^{\text{MC},x}(t) = \int_{0}^{+\infty} a^{\text{desired},x}(t) \cdot f(t-s) \, ds$$

841 Where:

$$f(t) = k_1 \cdot e^{-t/T_1} + k_2 \cdot e^{-t/T_2} + k_3 \cdot e^{-t/T_3},$$

842

 $T = \begin{bmatrix} .07 & .3 & 1 \end{bmatrix}, K = \begin{bmatrix} -0.4254 & 1.9938 & -0.5684 \end{bmatrix}$

844 These coefficients were adjusted to respect the following constraints:

845 -f(0) = 1, i.e. the output would correspond to the input at t = 0. This was chosen to ensure that the 846 high-frequency content of the motion would be rendered by translating the platform.

847 $-\int_0^\infty f = 0$: This was chosen to ensure that, if the input was an infinitely long acceleration, the motion of 848 the platform would stabilize to a point where the linear velocity was 0.

849 -df/dt = 0 at t = 0. This was chosen because tilt velocity of the platform is equal to -df/dt. Since 850 the tilt velocity at t<0 is zero, this constraint ensures that tilt velocity is continuous and prevents excessive 851 angular acceleration at t=0.

The same process is repeated for the y component of the acceleration. 852

Finally, the amount of tilt (θ , in degrees) is calculated based on the difference between the desired platform 854 motion and the deliverable motion:

855
$$\theta_{t+1}^{\text{MC},x} = \sin^{-1} \left(\frac{a_{t+1}^{\text{moog},x} - a_{t+1}^{\text{MC},x}}{g} \right)$$

$$\theta_{t+1}^{MC,x} = \sin^{-1} \left(\frac{a_{t+1}^{moog,x} - a_{t+1}^{MC,x}}{g} \right)$$

856

$$\theta_{t+1}^{MC,y} = \sin^{-1} \left(\frac{a_{t+1}^{moog,y} - a_{t+1}^{MC,y}}{g} \right)$$

857 Where $g = 9.81 \ m_{s^2}$.

858 STEP 6:

859 Afterwards, the tilt velocity and acceleration are being calculated:

860
$$\dot{\theta}_{t+1}^{\text{MC},x} = \frac{\theta_{t+1}^{\text{MC},x} - \theta_{t}^{\text{MC},x}}{dt}, \ \dot{\theta}_{t+1}^{\text{MC},y} = \frac{\theta_{t+1}^{\text{MC},y} - \theta_{t}^{\text{MC},y}}{dt}$$

861
$$\ddot{\theta}_{t+1}^{\mathrm{MC},x} = \frac{\dot{\theta}_{t+1}^{\mathrm{MC},x} - \dot{\theta}_{t}^{\mathrm{MC},x}}{dt}, \ \ddot{\theta}_{t+1}^{\mathrm{MC},y} = \frac{\dot{\theta}_{t+1}^{\mathrm{MC},y} - \dot{\theta}_{t}^{\mathrm{MC},y}}{dt}$$

862

In a next step, we compute the motion command that should be sent by the platform. Note that the platform is placed at a height *h* below the head. Therefore, tilting the platform by an angle θ induces a linear displacement of the head corresponding to $-h \cdot \theta \cdot \pi/_{180}$. Therefore, a linear displacement is added to the platform's motion to compensate for this. Next, we limit the platform's acceleration, velocity and position commands to ensure that they remain within the limit of the actuators. For this purpose, we define the following function $f_{\lambda,x_{max}}(x)$:

869
$$\begin{cases} if |x| \le \lambda \cdot x_{max}, & f_{\lambda, x_{max}}(x) = x \\ else if |x| \le (2 - \lambda) \cdot x_{max}, & f_{\lambda, x_{max}}(x) = x_{max} \cdot sign(x) \cdot [|x/x_{max}| - \frac{1}{4 \cdot (1 - \lambda)} \cdot (|x/x_{max}| - \lambda)^2] \\ if |x| > (2 - \lambda) \cdot x_{max}, & f_{\lambda, x_{max}}(x) = sign(x) \cdot x_{max} \end{cases}$$

870

This function is designed so that if the input x increases continuously, e.g. x(t) = t, then the output $f_{\lambda,x_{max}}(x(t))$ will be identical to x until x reaches a threshold $\lambda \cdot x_{max}$. After this, the output will decelerate continuously $(\frac{d f_{\lambda,x_{max}}(x(t))}{dt} = constant)$ until it stops at a value x_{max} . We fed the platform's acceleration, velocity and position command through this function, as follows:

875
$$a_{t+1}^{\text{moog},x} = f_{\lambda,a_{max}}(a_{t+1}^{\text{MC},x} + h \cdot \ddot{\theta}_{t+1}^{\text{MC},x} \cdot \pi/180)$$

876
$$v_{t+1}^{\text{moog},x} = f_{\lambda,v_{max}}(v_t^{\text{moog},x} + dt. a_{t+1}^{\text{moog},x})$$

877
$$x_{t+1}^{\text{moog},x} = f_{\lambda,x_{max}}(x_t^{\text{moog},x} + dt, x_{t+1}^{\text{moog},x})$$

878

The same operation takes place for the y component of the acceleration, as well as for the platform velocity and position. The process is repeated for the tilt command itself.

We set $\lambda = 0.75$ and $a_{max} = 4 m/s^2$, $v_{max} = 0.4 m/s$, $x_{max} = 0.23 m$, $\ddot{\theta}_{max} = 300 \circ/s$, $\dot{\theta}_{max} = 300 \circ/s$, $\dot{\theta}_{max} = 300 \circ/s$ and $\theta_{max} = 10^\circ$, slightly below the platform's and actuator physical limits. This ensured that the platform's motion matched exactly the motion cueing algorithm's output, as long as it stayed within 75% of the platform's range. Otherwise, the function *f* ensured a smooth trajectory and, as detailed in STEP 8 to 10, a feedback mechanism was used to update the participant position in the VR environment, so as to guarantee that visual motion always matched inertial motion.

887 **STEP 7:**

888 The motor commands for tilt and translation are being sent to the platform:

889
$$\left[x_{t+1}^{\text{moog}}, y_{t+1}^{\text{moog}}, \theta_{t+1}^{\text{moog},x}, \theta_{t+1}^{\text{moog},y}\right]$$

890 <u>STEP 8:</u>

Because of STEP 6, the total Gravito-Inertial Acceleration (GIA) of the platform may differ from what is commanded by the motion cueing algorithm. To detect and discrepancy, we computed the GIA provided by the platform:

894
$$v_{t+1}^{\text{actual},x} = \frac{\left(x_{t+1}^{\text{moog}} - x_t^{\text{moog}}\right)}{dt}, \quad v_{t+1}^{\text{actual},y} = \frac{\left(y_{t+1}^{\text{moog}} - y_t^{\text{moog}}\right)}{dt}$$

895

896
$$a_{t+1}^{\text{actual},x} = \frac{\left(v_{t+1}^{\text{actual},x} - v_{t}^{\text{actual},x}\right)}{dt}, \qquad a_{t+1}^{\text{actual},y} = \frac{\left(v_{t+1}^{\text{actual},y} - y_{v}^{\text{actual},y}\right)}{dt}$$

897
$$GIA_{t+1}^{\operatorname{actual},x} = a_{t+1}^{\operatorname{actual},x} + g \cdot \sin \theta_{t+1}^{\operatorname{moog},x} - h \cdot \ddot{\theta}_{t+1}^{\operatorname{moog},x} \cdot \pi/180$$

898
$$GIA_{t+1}^{\text{actual},y} = a_{t+1}^{\text{actual},y} + g \cdot \sin \theta_{t+1}^{\text{moog},y} - h \cdot \ddot{\theta}_{t+1}^{\text{moog},y} \cdot \pi/180$$

899

900 <u>STEP 9:</u>

901 We transform platform's GIA, into participant's reference frame:

902
$$GIA_{t+1}^{\operatorname{sub},x} = GIA_{t+1}^{\operatorname{actual},x} \cdot \cos \varphi_t^{\operatorname{moog}} + GIA_{t+1}^{\operatorname{actual},y} \cdot \sin \varphi_t^{\operatorname{moog}}$$

903
$$GIA_{t+1}^{\operatorname{sub},y} = -GIA_{t+1}^{\operatorname{actual},x} \cdot \sin \varphi_t^{\operatorname{moog}} + GIA_{t+1}^{\operatorname{actual},y} \cdot \cos \varphi_t^{\operatorname{moog}}$$

904

905 Also, the error e^{sub} between the obtained GIA and desired GIA (from STEP 3) is calculated, and fed 906 through the same sigmoid function ($\lambda = .75$, $GIA_{max} = 1 \frac{m}{s^2}$) discussed previously, to avoid 907 computational instability in the case of a big mismatch:

908
$$e_{t+1}^{x} = f_{\lambda,GIA_{max}}(GIA_{t+1}^{sub,x} - a_{t+1}^{sub,x})$$

909

$$e_{t+1}^{y} = f_{\lambda,GIA_{max}}(GIA_{t+1}^{\mathrm{sub},y} - a_{t+1}^{\mathrm{sub},y})$$

910 STEP 10:

911 The GIA error is now used to update the system in the case of a mismatch. First, it is transformed into VR 912 coordinates. Then, the velocity and position in VR coordinates are recomputed based on the joystick input 913 and on the error signal:

914
$$e_{t+1}^{\text{VR},x} = e_{t+1}^{\sup,x} \cdot \cos \varphi_t^{\text{VR}} - e_{t+1}^{\sup,y} \cdot \sin \varphi_t^{\text{VR}}$$

915
$$e_{t+1}^{\text{VR},y} = e_{t+1}^{\sup,x} \cdot \sin \varphi_t^{\text{VR}} + e_{t+1}^{\sup,y} \cdot \cos \varphi_t^{\text{VR}}$$

916
$$\hat{v}_{t+1}^{\text{VR},x} = \hat{v}_t^{\text{VR},x} + \left(a_{t+1}^{\text{VR},x} + e_{t+1}^{\text{VR},x}\right) \cdot dt$$

917
$$\hat{v}_{t+1}^{\text{VR},y} = \hat{v}_t^{\text{VR},y} + \left(a_{t+1}^{\text{VR},y} + e_{t+1}^{\text{VR},y}\right) \cdot dt$$

918
$$x_{t+1}^{\text{VR},x} = x_t^{\text{VR},x} + \hat{v}_{t+1}^{\text{VR},x} \cdot dt$$

918
919

$$x_{t+1}^{VR,y} = x_t^{VR,y} + \hat{v}_{t+1}^{VR,y} \cdot dt$$

919
 $x_{t+1}^{VR,y} = x_t^{VR,y} + \hat{v}_{t+1}^{VR,y} \cdot dt$

920
$$\varphi_{t+1}^{\text{VR}} = \varphi_t^{\text{VR}} + \omega_t^{\text{VR}} \cdot dt$$

921 Note that the error signal is also fed into the acceleration in VR coordinates (see STEP 3). Ideally, linear 922 acceleration should be computed based on the updated velocity value at time t i.e.:

923

924
$$\alpha_{t+1}^{\text{VR},x} = \frac{v_{t+1}^{\text{VR},x} - \hat{v}_t^{\text{VR},x}}{dt}$$

925 However, we found that this led to numerical instability, and instead we introduced a time constant τ_{MC} = 926 1s in the computation, as shown in STEP 3.

- 927
- 928

929 930 931	References	Alefantis, P., Lakshminarasimhan, K. J., Avila, E., Pitkow, X., & Angelaki, D. E. (2021). Sensory evidence accumulation using optic flow in a naturalistic navigation task. <i>Biorxiv</i> .
932 933 934		Angelaki, D. E., & Dickman, J. D. (2000). Spatiotemporal processing of linear accelerations: Primary afferent and central vestibular neuron responses. <i>Journal of Neurophysiology</i> . https://doi.org/10.1152/jn.2000.84.4.2113
935 936 937		Arce, F., Novick, I., Shahar, M., Link, Y., Ghez, C., & Vaadia, E. (2009). Differences in context and feedback result in different trajectories and adaptation strategies in reaching. <i>PLoS ONE</i> . https://doi.org/10.1371/journal.pone.0004214
938 939 940		Arthur, J. C., Kortte, K. B., Shelhamer, M., & Schubert, M. C. (2012). Linear path integration deficits in patients with abnormal vestibular afference. <i>Seeing and Perceiving</i> . https://doi.org/10.1163/187847612X629928
941 942 943		Batcho, C. S., Gagné, M., Bouyer, L. J., Roy, J. S., & Mercier, C. (2016). Impact of online visual feedback on motor acquisition and retention when learning to reach in a force field. <i>Neuroscience</i> . https://doi.org/10.1016/j.neuroscience.2016.09.020
944 945 946		Bergmann, J., Krauß, E., Münch, A., Jungmann, R., Oberfeld, D., & Hecht, H. (2011). Locomotor and verbal distance judgments in action and vista space. <i>Experimental Brain Research</i> . https://doi.org/10.1007/s00221-011-2597-z
947 948		Berniker, M., Voss, M., & Kording, K. (2010). Learning priors for bayesian computations in the nervous system. <i>PLoS ONE</i> . https://doi.org/10.1371/journal.pone.0012686
949 950 951		Burdet, E., Osu, R., Franklin, D. W., Milner, T. E., & Kawato, M. (2001). The central nervous system stabilizes unstable dynamics by learning optimal impedance. <i>Nature</i> . https://doi.org/10.1038/35106566
952 953		Campos, J. L., Butler, J. S., & Bülthoff, H. H. (2012). Multisensory integration in the estimation of walked distances. <i>Experimental Brain Research</i> . https://doi.org/10.1007/s00221-012-3048-1

954 955 956	Campos, J. L., Byrne, P., & Sun, H. J. (2010). The brain weights body-based cues higher than vision when estimating walked distances. <i>European Journal of Neuroscience</i> . https://doi.org/10.1111/j.1460-9568.2010.07212.x
957	Chen, X., McNamara, T. P., Kelly, J. W., & Wolbers, T. (2017). Cue combination in human spatial
958	navigation. <i>Cognitive Psychology</i> . https://doi.org/10.1016/j.cogpsych.2017.04.003
959	Chrastil, E. R., Nicora, G. L., & Huang, A. (2019). Vision and proprioception make equal
960	contributions to path integration in a novel homing task. <i>Cognition</i> .
961	https://doi.org/10.1016/j.cognition.2019.06.010
962	Chrastil, E. R., Sherrill, K. R., Hasselmo, M. E., & Stern, C. E. (2016). Which way and how far?
963	Tracking of translation and rotation information for human path integration. <i>Human Brain</i>
964	<i>Mapping</i> . https://doi.org/10.1002/hbm.23265
965 966	Einstein, A. (1907). Relativitätsprinzip und die aus demselben gezogenen Folgerungen [On the Relativity Principle and the Conclusions Drawn from It]. <i>Jahrbuch Der Radioaktivität</i> .
967	Fernandez, C., & Goldberg, J. M. (1976). Physiology of peripheral neurons innervating otolith organs
968	of the squirrel monkey. I. Response to static tilts and to long duration centrifugal force. <i>Journal</i>
969	of Neurophysiology. https://doi.org/10.1152/jn.1976.39.5.970
970	Fernandez, C., Goldberg, J. M., & Abend, W. K. (1972). Response to static tilts of peripheral neurons
971	innervating otolith organs of the squirrel monkey. <i>Journal of Neurophysiology</i> .
972	https://doi.org/10.1152/jn.1972.35.6.978
973	Glasauer, S, Amorim, M. A., Vitte, E., & Berthoz, A. (1994). Goal-directed linear locomotion in
974	normal and labyrinthine-defective subjects. <i>Experimental Brain Research</i> .
975	https://doi.org/10.1007/BF00228420
976	Glasauer, Stefan, Schneider, E., Grasso, R., & Ivanenko, Y. P. (2007). Space-time relativity in self-
977	motion reproduction. <i>Journal of Neurophysiology</i> . https://doi.org/10.1152/jn.01243.2005
978 979 980	Grasso, R., Glasauer, S., Georges-François, P., & Israël, I. (1999). Replication of passive whole-body linear displacements from inertial cues. <i>Annals of the New York Academy of Sciences</i> . https://doi.org/10.1111/j.1749-6632.1999.tb09197.x
981	Israël, I., Bronstein, A. M., Kanayama, R., Faldon, M., & Gresty, M. A. (1996). Visual and vestibular
982	factors influencing vestibular "navigation." <i>Experimental Brain Research</i> .
983	https://doi.org/10.1007/BF00227947
984 985 986	Israël, I., Grasso, R., Georges-François, P., Tsuzuku, T., & Berthoz, A. (1997). Spatial memory and path integration studied by self-driven passive linear displacement. I. Basic properties. <i>Journal of Neurophysiology</i> . https://doi.org/10.1152/jn.1997.77.6.3180
987	Jürgens, R., & Becker, W. (2006). Perception of angular displacement without landmarks: Evidence
988	for Bayesian fusion of vestibular, optokinetic, podokinesthetic, and cognitive information.
989	<i>Experimental Brain Research</i> . https://doi.org/10.1007/s00221-006-0486-7
990	Karmali, F., Lim, K., & Merfeld, D. M. (2014). Visual and vestibular perceptual thresholds each
991	demonstrate better precision at specific frequencies and also exhibit optimal integration.
992	<i>Journal of Neurophysiology</i> . https://doi.org/10.1152/jn.00332.2013
993	Kearns, M. J., Warren, W. H., Duchon, A. P., & Tarr, M. J. (2002). Path integration from optic flow
994	and body senses in a homing task. <i>Perception</i> . https://doi.org/10.1068/p3311
995	Kingma, H. (2005). Thresholds for perception of direction of linear acceleration as a possible
996	evaluation of the otolith function. <i>BMC Ear, Nose and Throat Disorders</i> .
997	https://doi.org/10.1186/1472-6815-5-5

998 999	Klatzky, R. L., Loomis, J. M., Beall, A. C., Chance, S., & Golledge, R. G. (1998). Spatial updating of selfposition and orientation during real. <i>Psychological Science</i> .
1000	Koppen, M., Horst, A. C. T., & Pieter Medendorp, W. (2019). Weighted Visual and Vestibular Cues
1001	for Spatial Updating during Passive Self-Motion. <i>Multisensory Research</i> .
1002	https://doi.org/10.1163/22134808-20191364
1003 1004	Körding, K. P., Ku, S. P., & Wolpert, D. M. (2004). Bayesian integration in force estimation. <i>Journal of Neurophysiology</i> . https://doi.org/10.1152/jn.00275.2004
1005	Kording, K. P., Tenenbaum, J. B., & Shadmehr, R. (2007). The dynamics of memory as a
1006	consequence of optimal adaptation to a changing body. <i>Nature Neuroscience</i> .
1007	https://doi.org/10.1038/nn1901
1008	Krakauer, J. W., Ghilardi, M. F., & Ghez, C. (1999). Independent learning of internal models for
1009	kinematic and dynamic control of reaching. <i>Nature Neuroscience</i> .
1010	https://doi.org/10.1038/14826
1011	Lackner, J. R., & Dizio, P. (1994). Rapid adaptation to Coriolis force perturbations of arm trajectory.
1012	Journal of Neurophysiology. https://doi.org/10.1152/jn.1994.72.1.299
1013	Lakshminarasimhan, K. J., Petsalis, M., Park, H., DeAngelis, G. C., Pitkow, X., & Angelaki, D. E.
1014	(2018). A Dynamic Bayesian Observer Model Reveals Origins of Bias in Visual Path
1015	Integration. <i>Neuron</i> . https://doi.org/10.1016/j.neuron.2018.05.040
1016	Lim, K., Karmali, F., Nicoucar, K., & Merfeld, D. M. (2017). Perceptual precision of passive body
1017	tilt is consistent with statistically optimal cue integration. <i>Journal of Neurophysiology</i> .
1018	https://doi.org/10.1152/jn.00073.2016
1019	MacNeilage, P. R., Turner, A. H., & Angelaki, D. E. (2010). Canal-otolith interactions and detection
1020	thresholds of linear and angular components during curved-path self-motion. <i>Journal of</i>
1021	<i>Neurophysiology</i> . https://doi.org/10.1152/jn.01067.2009
1022	Mathys, C., Daunizeau, J., Friston, K. J., & Stephan, K. E. (2011). A Bayesian foundation for
1023	individual learning under uncertainty. <i>Frontiers in Human Neuroscience</i> .
1024	https://doi.org/10.3389/fnhum.2011.00039
1025	Péruch, P., Borel, L., Gaunet, F., Thinus-Blanc, G., Magnan, J., & Lacour, M. (1999). Spatial
1026	performance of unilateral vestibular defective patients: In nonvisual versus visual navigation.
1027	<i>Journal of Vestibular Research: Equilibrium and Orientation</i> .
1028	Péruch, P., Borel, L., Magnan, J., & Lacour, M. (2005). Direction and distance deficits in path
1029	integration after unilateral vestibular loss depend on task complexity. <i>Cognitive Brain</i>
1030	<i>Research</i> . https://doi.org/10.1016/j.cogbrainres.2005.09.012
1031	Petrini, K., Caradonna, A., Foster, C., Burgess, N., & Nardini, M. (2016). How vision and self-
1032	motion combine or compete during path reproduction changes with age. <i>Scientific Reports</i> .
1033	https://doi.org/10.1038/srep29163
1034	Petzschner, F. H., & Glasauer, S. (2011). Iterative Bayesian estimation as an explanation for range
1035	and regression effects: A study on human path integration. <i>Journal of Neuroscience</i> .
1036	https://doi.org/10.1523/JNEUROSCI.2028-11.2011
1037	Prsa, M., Jimenez-Rezende, D., & Blanke, O. (2015). Inference of perceptual priors from path
1038	dynamics of passive self-motion. <i>Journal of Neurophysiology</i> .
1039	https://doi.org/10.1152/jn.00755.2014
1040	Sağlam, M., Glasauer, S., & Lehnen, N. (2014). Vestibular and cerebellar contribution to gaze
1041	optimality. <i>Brain</i> . https://doi.org/10.1093/brain/awu006

1042	Seemungal, B. M., Glasauer, S., Gresty, M. A., & Bronstein, A. M. (2007). Vestibular perception and
1043	navigation in the congenitally blind. <i>Journal of Neurophysiology</i> .
1044	https://doi.org/10.1152/jn.01321.2006
1045 1046	Seidman, S. H. (2008). Translational motion perception and vestiboocular responses in the absence of non-inertial cues. <i>Experimental Brain Research</i> . https://doi.org/10.1007/s00221-007-1072-3
1047 1048	Shadmehr, R., & Mussa-Ivaldi, F. A. (1994). Adaptive representation of dynamics during learning of a motor task. <i>Journal of Neuroscience</i> . https://doi.org/10.1523/jneurosci.14-05-03208.1994
1049	Stangl, M., Kanitscheider, I., Riemer, M., Fiete, I., & Wolbers, T. (2020). Sources of path integration
1050	error in young and aging humans. <i>Nature Communications</i> . https://doi.org/10.1038/s41467-
1051	020-15805-9
1052	Takahashi, C. D., Scheidt, R. A., & Reinkensmeyer, D. J. (2001). Impedance control and internal
1053	model formation when reaching in a randomly varying dynamical environment. <i>Journal of</i>
1054	<i>Neurophysiology</i> . https://doi.org/10.1152/jn.2001.86.2.1047
1055	Tcheang, L., Bülthoff, H. H., & Burgess, N. (2011). Visual influence on path integration in darkness
1056	indicates a multimodal representation of large-scale space. <i>Proceedings of the National</i>
1057	<i>Academy of Sciences of the United States of America</i> . https://doi.org/10.1073/pnas.1011843108
1058	ter Horst, A. C., Koppen, M., Selen, L. P. J., & Pieter Medendorp, W. (2015). Reliability-based
1059	weighting of visual and vestibular cues in displacement estimation. <i>PLoS ONE</i> .
1060	https://doi.org/10.1371/journal.pone.0145015
1061	Tramper, J. J., & Medendorp, W. P. (2015). Parallel updating and weighting of multiple spatial maps
1062	for visual stability during whole body motion. <i>Journal of Neurophysiology</i> .
1063	https://doi.org/10.1152/jn.00576.2015
1064	Zupan, L. H., & Merfeld, D. M. (2008). Interaural self-motion linear velocity thresholds are shifted
1065	by roll vection. <i>Experimental Brain Research</i> . https://doi.org/10.1007/s00221-008-1540-4
1066	
1067	