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2 3 4	Oculomotor plant and neural dynamics suggest gaze control requires integration on distributed timescales			
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30	Running head: Response tir	nescales in the zebrafish ocul	omotor plant	
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KEY POINTS

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36 37 38 39	• Recent observations of oculomotor plant response properties and neural a across the oculomotor system have called into question classical formulations the oculomotor plant and the oculomotor integrator.	•
40 41 42 43	• Here we use measurements from new and published experiments in the zebrafish together with modelling to reconcile recent oculomotor plant obser with oculomotor integrator function.	
44 45 46 47	• We developed computational techniques to characterize oculomotor plant resover several seconds in awake animals, demonstrating that long timescale resover in anesthetized animals extend to the awake state.	-
48 49 50 51	• Analysis of firing patterns of oculomotor integrator neurons demonstra sufficiency of this activity for stabilizing gaze given an oculomotor plan multiple, distributed response timescales.	
52 53 54	• Our results support a formulation of gaze stabilization by the oculomotor symbic commands for stabilizing gaze are generated through integration on m distributed timescales.	

55 ABSTRACT

56 A fundamental principle of biological motor control is that the neural commands driving movement must conform to the response properties of the motor plants they control. In the 57 58 oculomotor system, characterizations of oculomotor plant dynamics traditionally supported 59 models in which the plant responds to neural drive to extraocular muscles on exclusively 60 short, subsecond timescales. These models predict that the stabilization of gaze during 61 fixations between saccades requires neural drive that approximates eye position on longer 62 timescales and is generated through the temporal integration of brief eye velocity-encoding 63 signals that cause saccades. However, recent measurements of oculomotor plant behaviour 64 have revealed responses on longer timescales, and measurements of firing patterns in the 65 oculomotor integrator have revealed a more complex encoding of eye movement dynamics. 66 Here we use measurements from new and published experiments in the larval zebrafish to 67 link dynamics in the oculomotor plant to dynamics in the neural integrator. The oculomotor 68 plant in both anaesthetized and awake larval zebrafish was characterized by a broad distribution of response timescales, including those much longer than one second. Analysis 69 70 of the firing patterns of oculomotor integrator neurons, which exhibited a broadly distributed 71 range of decay time constants, demonstrates the sufficiency of this activity for stabilizing 72 gaze given an oculomotor plant with distributed response timescales. This work suggests that 73 leaky integration on multiple, distributed timescales by the oculomotor integrator reflects an 74 inverse model for generating oculomotor commands, and that multi-timescale dynamics may be a general feature of motor circuitry. 75

76 **INTRODUCTION**

77 Motor plants transform commands from motor neurons into action. Because motor plant 78 responses to neural drive can be complex and history-dependent, commands needed to elicit 79 a particular movement differ from the desired patterns of muscle activation. Standard models 80 of motor control assume that premotor circuitry generates appropriate motor commands by 81 filtering intended muscle activation through an inverse model of the plant being controlled 82 (Kawato 1999; Lisberger 2009). This filtering is then cancelled by the plant's response to the 83 command, resulting in the intended movement. In this manner, the filtering via the inverse 84 model compensates for the response properties of the plant.

85 This inverse model framework has proven useful in understanding motor command generation in the oculomotor system (Figure 1A; Green et al. 2007; Robinson 1989; Van 86 87 Opstal et al. 1985). In the classical view, based on the work of Robinson (1964), passive 88 oculomotor plant behaviour in the horizontal plane is modelled by a pair of one-dimensional 89 viscoelastic (Voigt) elements in series, each characterized by a time constant that dictates the 90 exponential time course of the element's length change following a step change in applied 91 force. These time constants have been estimated to be relatively short: 10-60 ms and 250-92 660 ms (Goldstein 1984; Optican and Miles 1985; Robinson 1964; Sklavos et al. 2006; Stahl 93 and Simpson 1995; Stahl et al. 2015). The motor command required for step changes in eye 94 position (saccades), determined by inverting this two-element plant model, is composed of 95 three components, each of which compensates for a different aspect of the plant model's 96 response properties (Goldstein 1984; Optican and Miles 1985; Robinson 1964). The first 97 component is a brief eye velocity-encoding burst of firing (termed the "pulse") that 98 overcomes plant viscosity to quickly pull the eye to a new position. The second is an eye 99 position-encoding "step" that counters the plant's elasticity to maintain the eye at a fixed 100 position during fixation. The final component is an exponential decay (termed the "slide") 101 with a time constant intermediate between the viscoelastic element time constants (see 102 Mathematical Appendix), which reflects the attenuating force needed to stabilize gaze as the 103 viscoelastic elements equilibrate following the saccade.

Each component appears to be reflected in the firing patterns of ocular motor neuronsduring horizontal eye movements (Robinson 1981). The pulse arises from saccadic burst

106 neurons that project to the ocular motor nuclei. The step (Cohen and Komatsuzaki 1972; 107 Robinson 1989; Scudder et al. 2002; Skavenski and Robinson 1973) and slide (Aksay et al. 108 2000; McFarland and Fuchs 1992) components have been observed in the firing of premotor 109 neurons constituting the velocity-to-position neural integrator for horizontal eye movements 110 (hVPNI), which receives eye velocity burst signals and appears to compute their temporal 111 integral, producing the step (Figure 1); the origins of the slide are less clear. Since gaze 112 stability in the dark, when the oculomotor system cannot rely on visual feedback to generate 113 motor commands, far exceeds that which could be attributed to the oculomotor plant alone, 114 premotor circuits appear to use an inverse plant model that enables substantial gaze stability 115 without visual feedback.

116 More recent work has exposed deficiencies in the classical view of eye movement 117 command generation. Sklavos et al. (2006; 2005) found evidence of additional time constants 118 on the order of 1 and 10 s following long steps of force externally applied to the eye. Quaia 119 et al. (2009) analysed the response of primate extraocular muscle to elongation steps, 120 demonstrating muscle tension relaxation on a wide range of timescales, with time constants 121 ranging up to at least 40 s. Davis-Lopez de Carrizosa et al. (2011) measured lateral rectus 122 muscle tension in cats, finding that it decays on timescales of 1 to 10 s during fixations 123 between saccades while eye position is approximately stable. Additionally, firing rates of 124 abducens motor neurons in primates during different types of eye movement are not 125 consistent with a common two-element plant model (Sylvestre and Cullen 1999). These 126 results support an expanded model of the oculomotor plant having several viscoelastic 127 elements with time constants distributed across several orders of magnitude, from 100 ms to 128 10 s (Sklavos et al. 2005, 2006). The long timescale responses of such plant models imply 129 that additional drive components that decay on long timescales are necessary to produce 130 stable fixations (Figure 1B,C; Sklavos et al. 2005).

However, questions remain about the relevance of long response timescales in the awake, behaving (active) state. Previous measurements of oculomotor plant dynamics in alert animals have been limited to short time intervals (< 400 ms in monkey, Anderson et al. 2009; < 2 s in mouse, Stahl et al. 2015), precluding observation of long timescale responses. These studies also used brief eye position steps < 1 s, which would not appreciably deform 136 viscoelastic elements with long time constants (Anderson et al. 2009; Sklavos et al. 2006).

- Furthermore, previous fitting of oculomotor plant models has depended on the assumption
 that viscoelastic elements were at equilibrium (Sklavos et al., 2005) and was therefore not
- 139 suitable for fitting model parameters in awake animals.

140 Other recent observations have revealed that neural activity in the hVPNI and ocular 141 motor neurons does not simply encode eye position and a single fast slide, as predicted by 142 classical models. We have reported that during fixations, firing rates in hVPNI neurons decay 143 on long timescales that vary across an order of magnitude within individual larval zebrafish 144 (Miri et al. 2011a). Such heterogeneity in firing rate decay timescales has also been measured 145 during fixation in adult goldfish hVPNI (Miri et al. 2011a) and in monkey oculomotor 146 integrator neurons (Joshua et al. 2013). In cats, abducens firing after saccades decays on 147 varying timescales generally greater than 1 s, with such decays believed to arise from the 148 oculomotor integrator (Davis-Lopez de Carrizosa et al. 2011). However, it remains to be seen 149 whether firing in the hVPNI could constitute a signal sufficient to stabilize a plant with 150 distributed response timescales.

151 Here we used measurements of oculomotor plant dynamics and analysis of previously 152 obtained neural recordings in the larval zebrafish to assess whether the hVPNI could be 153 implementing an approximate inverse model that accommodates long timescale behaviour of 154 the oculomotor plant to promote gaze stability. We performed very long mechanical 155 displacements in both anaesthetized and awake, behaving animals in order to deform long 156 timescale response elements. We developed analytical methods for measuring the eye's 157 return from long displacements in awake, behaving animals. Our measurements demonstrate 158 the existence of both short (< 1 s time constant) and long (> 1 s) response timescales in the 159 larval zebrafish oculomotor plant in both anaesthetized and awake animals. We used these 160 measurements to fit oculomotor plant models for each animal, employing a new method that 161 does not require an equilibrium assumption. We then compared the predictions of the neural 162 drive during active state fixations with measurements of activity in the larval zebrafish 163 hVPNI. Analysis of the distribution of decay times seen in hVPNI neuron firing rates suggests that this distribution is sufficient to enable stabilization of an oculomotor plant with 164 165 the distributed response timescales we observed, including those longer than 1 s. Our results

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- 166 support a view of integration in the oculomotor system in which hVPNI firing, rather than
- 167 purely or primarily encoding eye position, compensates for plant viscoelasticity by
- 168 generating firing rate decay on multiple, distributed timescales (Figure 1).

169 METHODS

170 Ethical approval

All experiments were performed in compliance with protocols approved by the Princeton University Institutional Animal Care and Use Committee (protocols #1726 and 1863), and in accordance with the policies of the *Journal of Physiology*. Following these guidelines ensured that animal distress was minimized in the course of our study.

175

189

176 Framework for modelling oculomotor plant

We modelled the oculomotor plant as a combination of viscoelastic Voigt elements in series
that respond over *n* effective timescales. This was represented as a linear filter whose impulse
response function consists of a sum of exponentially decaying components (Robinson 1964;
Sklavos et al. 2005),

181
$$p(t) = \sum_{i=1}^{n} c_i e^{-t/\tau_i},$$
 (1)

182 where c_i is the coefficient of the component with time constant τ_i . The time constants can be 183 identified by finding the step response of the system. If a force f(t) is applied to the system 184 at time t = 0, the measured eye position y(t) will be a convolution of the force profile and 185 the impulse response,

186
$$y(t) = (f * p)(t) = \int_0^t du \ f(u) p(t-u).$$
(2)

187 If the applied force is stopped at time $t = t_0$, then the measured eye position at later times 188 will be

$$y(t > t_0) = \int_0^{t_0} du f(u) \sum_{i=1}^n c_i e^{-(t-u)/\tau_i}$$

= $\sum_{i=1}^n \left(c_i \int_0^{t_0} du f(u) e^{-(t_0-u)/\tau_i} \right) e^{-(t-t_0)/\tau_i}$
= $\sum_{i=1}^n a_i e^{-(t-t_0)/\tau_i}$, (3)

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190 that is, the response after release will be a sum of exponentials with the same time constants

as the plant.

Based on the above, in order to model the oculomotor plant, we took the followingsteps, detailed in the sections below:

- 194 1. Apply a transient external force ("displacement") resulting in a step change in eye 195 position, and measure the eye position after release from displacement ("step 196 response").
- 197 2. Fit a multiexponential model to the step response and extract the plant time constants 198 $\{\tau_i\}$.
- 199 3. Find the plant coefficients $\{c_i\}$ by fitting a model of the form in equation (2) to the 200 eye position during and after displacement.
- 201

202 Step response measurement

mitfa^{-/-} (nacre) mutant zebrafish (*Danio rerio*) larvae (Lister et al. 1999) ages 5-8 days postfertilization were used for all experiments. We obtained the nacre strain from Zebrafish
International Resource Center. Embryos were reared in egg water (Westerfield, 2007) in petri
dishes in an incubator at 28°C on a 12 h light/12 h dark cycle. Larvae at this age feed from
their yolk and additional food was not provided.

To enable eye tracking, larvae were immobilised by embedding in a thin layer of 1.7% low melting point agarose (SeaPlaque, Lonza) immediately prior to data collection, and the agarose was removed from around the eyes to allow free eye movement. A rectangular agarose block containing the larva was excised and mounted on a Sylgard platform in a waterfilled chamber. Individual larvae were embedded for no more than 3 hours. Following data collection, larvae were removed from agarose and immediately euthanized by submerging in ice water for > 5 minutes, to which bleach was added to a concentration of 1% by volume.

We used one of two methods to anaesthetize larvae. Ethyl 3-aminobenzoate methanesulfonate (MS-222, Sigma; n = 10 larvae) was gradually added to the chamber water to achieve a concentration at which spontaneous eye movements stopped. Final concentrations were between 0.005 and 0.015% (weight/volume). For ketamine experiments (n = 6 larvae), embedded larvae were incubated for 15 minutes in 0.5% (weight/volume) ketamine prior to mounting in the chamber; no ketamine was added to the chamber water.
This ketamine concentration was found to be sufficient to abolish spontaneous eye movement
in most larvae. Data were not collected from larvae that performed eye movements under
anaesthesia. Separate sets of larvae were used for each of the experimental groups: MS-222
anesthetized, ketamine anesthetized, and awake.

225 The left eye's response to step displacement was measured in the dark. In order to 226 displace the eye, a blunt probe (hemispherical tip $\sim 30 \ \mu m$ in diameter) controlled by a 227 hydraulic micromanipulator (Siskiyou) was brought toward a point ~50 µm temporal of the 228 centre of the left eye at an oblique angle relative to the eye's minor axis (considering the eye 229 as an ellipsoid; Figure 2A). After gently contacting the eye, the probe was advanced to rotate 230 the eye temporally in the horizontal plane. Probe advancement was performed quickly, taking 231 less than a second. For active state measurements (n = 6 larvae), this displacement was 232 performed 5-7 s following a saccade in which the eye moved nasally. This allowed the 233 expected position of the eye in the absence of the displacement to be estimated by 234 extrapolating a fit to the eye position between the saccade and the displacement (see below). 235 The eye was released by quickly retracting the probe. Eye position was tracked at ~ 70 Hz for 236 > 60 seconds following release using methods previously described (Beck et al. 2004; Miri 237 et al. 2011b). Briefly, a 945-nm LED illuminated the chamber from below while a mirror, 238 long-pass filter, and charge-coupled device (CCD) camera above the chamber collected video 239 images that were processed in real-time to extract eye position measurements using software 240 custom written in LabView (National Instruments). In this software, two regions of interest 241 (ROIs) that included either the eyes or a fixed segment of the body were drawn on a reference 242 CCD image. During data collection, the ROIs were thresholded, the two largest objects 243 within the eye ROI were defined as the eyes, the largest object within the body field was 244 defined as the body segment, and the edges of these three objects were smoothed. The body 245 axis was defined as the line connecting the centre of the body segment and the midpoint 246 between the centroids of the eyes. Horizontal eye positions were measured as the angle 247 formed by the major axis of the eye and the body axis. Eye position measurements were 248 digitized by a Digidata 1440A (Molecular Devices) and recorded at 5000 Hz in Clampex 249 (v.10, Molecular Devices). Visual inspection of eye tracking images indicated that eye shape was at most minimally disturbed by contact with the probe and any disturbance was confined to the site of probe contact. There was no visible deformation of the eye from contact with the probe that extended appreciably into the displacement response.

In each anaesthetized larva, two displacements of differing duration were performed: 253 254 10 and 60 s in MS-222 experiments, 15 and 90 s in ketamine experiments. Displacements ranged from 14.8 to 22.2° under MS-222, and 8.5 to 20.9° under ketamine. Pairs of 255 256 displacements performed on individual larvae were nearly equal, differing on average by 257 only 1.5°. In awake larvae, up to five displacements were performed on each larva, each 258 between 6.5 and 8.5 s in duration, and 11.6 and 26.0° in size. Active state trials in which 259 spontaneous eye movements occurred during the applied displacement or within 8 s 260 following the release were discarded. As a result, at most two responses from each larva were 261 analysed. Spontaneous eye movements during displacement could be identified by motion of 262 the undisplaced eye. At least 10 minutes elapsed between displacements.

263

264 Step response fitting

Eye position during the displacement prior to release was measured from an image captured while the eye was displaced. The time of release was defined as the time of the last sample during which the probe was contiguous with the eye in the video image. For MS-222 and ketamine experiments, baseline eye position was measured as the mean eye position during a 50 second epoch preceding the displacement and was subtracted from the eye position time series.

271 For active state responses, the centre of gaze was estimated from a plot of eye velocity 272 versus eye position (a "PV plot"; Becker and Klein 1973; Goldman et al. 2002) assembled as 273 follows from at least three minutes of eye position data collected during spontaneous eye 274 movement prior to the displacement. First, data from 100 ms prior to each saccade until 500 275 ms after each saccade were discarded. The remaining time series data were divided into non-276 overlapping 0.3 s segments. Next, the mean eye position and the slope of a least-squares fit 277 line to eye position over each segment were calculated to define the two coordinates of points 278 comprising the PV plot. Finally, a linear function was least-squares fit to these points. The intercept of this function with the eye position axis was defined as the centre of gaze andsubtracted from all measurements in the eye position time series.

281 Since data were initially acquired at ~70 Hz and digitized at 5000 Hz, we 282 downsampled eye position time series before performing any data analysis. We therefore 283 subsampled traces every 72 time points, resulting in new traces at 69.44 Hz that we then 284 analysed. Subsequent analysis of the power spectra of eye position traces showed 285 anomalously large peaks at ~30 Hz, which are likely artefacts. These were removed, while 286 preserving the phase in each frequency bin, by scaling the amplitudes of the Fourier transform 287 in the peaks so that the amplitude was equal to that of the mean amplitude of the 6 frequency 288 bins surrounding the peak (3 closest on each side of the peak). All analyses were performed 289 in Python 3.7, using the Scientific Python stack (SciPy and NumPy).

290

Anaesthetized step responses Eye position step responses (Figure 2B) in anaesthetized larvae
 were fit with a multiexponential model,

293
$$y(t) = \sum_{i=1}^{n} a_i e^{-k_i t} + \varepsilon(t),$$
 (4)

where ε is independent Gaussian noise with mean 0 and variance σ^2 , $k_i = 1/\tau_i$ are inverse time constants, and the number of components *n* ranged between 1 and 6. Each coefficient was constrained to be nonnegative, $a_i \ge 0$, and the sum of the coefficients was constrained to equal 1. Visual inspection of these time series near the release time found ringing/oscillation present during the initial 50-200 ms following release in some cases, perhaps resulting from the manual control of the hydraulic manipulator. We therefore analysed responses beginning 230 ms and ending 60 s after release time

For each larva, we simultaneously fit the short and long step response with models of the form in equation (4) for each value of *n* between 1 and 6. For each value of *n*, we defined the best *n*-component model to be the one which maximized the sum \mathcal{L} of log-likelihoods \mathcal{L}_j for each response *j*,

$$\mathcal{L} = \sum_{j} \mathcal{L}_{j}.$$
 (5)

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306 Here,

307
$$\mathcal{L}_{j} = -\frac{T}{2}\log\frac{2\pi}{\psi_{j}} - \frac{\psi_{j}}{2}\sum_{k=1}^{T}(y(t_{k}) - \hat{y}(t_{k}))^{2}, \qquad (6)$$

308 where T is the total number of time points $\{t_k\}$ recorded in the response, y_i is the recorded eye position, \hat{y}_i is a model in the form of equation (4), and, for numerical stability, we defined 309 $\psi_i = 1/\sigma_i^2$ where σ_i is the standard deviation of the noise. We allowed the fits to each of the 310 two response durations to have different sets of coefficients $\{a_i\}$, but we required a single set 311 312 of inverse time constants $\{k_i\}$, with time constants constrained to be greater than 43.2 ms, or 313 3 samples. Separate standard deviations of the noise σ_i were fit for each response. To find 314 sets of parameters that maximized \mathcal{L} , we used the nonlinear solver Truncated Newton Conjugate-Gradient (TNC; implemented in the "optimize" library of SciPy), which we 315 provided with an analytical formula for the gradient of \mathcal{L} that was derived by hand. We used 316 100 initial sets of parameter values by choosing coefficients uniformly at random between 0 317 318 and 1 and then dividing each coefficient by the sum of all coefficients. Initial time constants 319 were chosen by taking random powers of 10, generated by first taking n evenly spaced 320 powers between -1 and 2, and adding Gaussian noise with mean 0 and standard deviation 0.1 to each. By examining the mean squared error curves of fits as n increased, we saw clear 321 322 "elbows" after which fit quality stopped visibly improving. Separately for each larva, we 323 called the value of n at which this elbow occurred n^* . For the sake of parsimony, we picked 324 the best overall model for each larva to be the best n^* -component model.

To examine the sensitivity of the parameter estimates, we used a parametric bootstrap procedure as follows. For each larva, we used the best step response fit (coefficients, time constants, and measurement noise variances) to generate 100 new eye position traces according to the model in equation (4), and re-ran our fitting procedure on each of these, then calculated the standard deviations of the resulting bootstrap parameters, which we used as an estimate of the standard deviations of the true parameter distributions. bioRxiv preprint doi: https://doi.org/10.1101/2021.06.30.450653; this version posted July 1, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

In order to determine the necessity of including long time constant components, we next repeated the above procedure to find the best fits for each larva when time constants were constrained to be less than 10 s.

334

335 Active state step responses Because the eye moves spontaneously in the active state and is 336 not at the centre of gaze prior to the imposed displacement, we did not model the active state 337 step response as a decay toward the centre of gaze. Rather, we modelled it as returning to 338 where the eye would have been had the displacement not occurred (Figure 2C). Let $y_{target}(t)$ 339 be the position of the eye if no displacement had occurred. Then, we modelled the active state 340 step response as

341
$$y_{\rm sr}(t - t_{\rm release}) = y(t) - y_{\rm target}(t), \tag{7}$$

342 where, as before, y(t) is the measured eye position.

We determined $y_{target}(t)$ by extrapolating the changing position of the eye, based on the 4-6 s of post-saccadic eye relaxation immediately preceding displacement (Figure 2C). We found that this relaxation could also be modelled well by equation (4), which we fit by maximum likelihood estimation to the eye position from 500 ms after the previous saccade to displacement onset, using the method described above for anaesthetized step responses. For all active state responses we picked a 2-component model for the best extrapolation, as there was negligible improvement in fit quality for >2 components.

To test the quality of this extrapolation technique, we identified long nasal fixations lasting at least 14.75 s in eye position recordings prior to each active step response. We then performed the same fitting procedure used to find $y_{target}(t)$ above, here fitting eye position from 500 ms to 6 s post-saccade (Figure 2D). We then calculated how much the resulting fit function deviated from the true eye position in a ~230 ms window of time centred at 14.64 s post-saccade by calculating the error relative to the extrapolation at each data point in this interval,

357 relative error at time
$$t = \frac{y(t) - y_{\text{target}}(t)}{y_{\text{target}}(t)}$$
. (8)

358 For each fixation, we then calculated the average relative error over the 16 data points in this 359 window (Figure 2D, inset; black point indicates the centre point of this window at time $t_{\text{extrap}} = 14.64$ s). A negative relative error meant that the extrapolation decayed more slowly 360 than the true fixation. We found that the average relative error was positive, so that fits were 361 362 likely to decay more quickly than the true fixation, but with relatively large variance (Figure 363 2D, right; mean \pm standard deviation [sd] = 0.14 \pm 0.32; n = 9). 3 out of the 9 fixations 364 resulted in a negative average relative error, where the extrapolated eye position decayed 365 more slowly than true eye position. Thus, a step response estimated from these extrapolations 366 would also decay more slowly than the "true" step response and would be more likely to have 367 long time constants. In order to make our fitting procedure more robust to such an artefact, 368 which could lead to spurious identification of long time constant decays, we sought to also fit a more quickly decaying, "conservative" extrapolation of eye position for each active state 369 370 step response.

371 We fit the conservative extrapolations by finding fit functions which fit the first 4-6 372 s of eye position well, but also decayed faster than the best extrapolation. We did this by 373 including a penalty term in our optimization that incentivized fit functions to pass through a 374 virtual point at time t_{extrap} that was closer to the null eye position than the best extrapolation 375 would be. First, we defined Δ to be the most negative relative error, averaged over the 376 window described above (Figure 2D, right, bottom blue point). Then, starting with the best 377 extrapolation (Figure 2C,D, red curve), we fit a higher order model, i.e., n + 1 components 378 if the best extrapolation needed *n* components, by maximum likelihood estimation, but where we augmented the model (4) by fitting an additional point $y(t_{extrap}) = (1 + \Delta)y_{target}(t_{extrap})$. 379 380 All of the fixations before displacement were less than 7 s long, so this additional point did not overwrite any actual data points. We let the noise at this additional point, $\varepsilon(t_{extrap})$, be 381 normally distributed with mean 0 and variance σ^2/λ . Increasing λ increasingly penalizes fits 382 that do not pass through $y(t_{extrap})$. By bisection search on λ , we allowed the procedure to 383 384 choose a fit whose sum of squared errors over the real data deviated from that of the best 385 extrapolation by ~10% as follows (Figure 2C,D, black curve). For each value of λ , we 386 performed maximum likelihood estimation from 100 starting points and picked the most parsimonious model, as described above (Anaesthetized step responses), to perform this comparison. The bisection search terminated when the deviation of the sum of squared errors fell between 9.9 and 10.1%. The eye position predicted by the maximum likelihood model for the value of λ for which the bisection search terminates was defined to be the conservative extrapolation.

392 For both the best extrapolation and conservative extrapolation, we calculated the 393 corresponding step response time series $y_{sr}(t)$. For each displacement in each larva, we fit 394 multiexponential models to both of these resulting active step response time series from 230 395 ms to between 8 and 14 s after release from displacement, with the fits performed as above 396 for anaesthetized responses. After discarding responses due to spontaneous eye movements, 397 a single response was fit for 3 of 6 larvae and a pair of responses was fit for the remaining 3. 398 We again ran the parametric bootstrap procedure, as in the anaesthetized case above, here 399 using the best three-component fits in order to evaluate the sensitivity of parameter estimates. 400 In order to evaluate the necessity of long time constant components, we also fit responses 401 while constraining the time constants to be less than 5 s. This was again done assuming the 402 best and conservative extrapolations.

For comparison of anaesthetized and active state results, we also fit three-component models as above to just the first 15 s of eye position following the 10 s displacement in larvae anaesthetized with MS-222.

406

407 **Oculomotor plant model estimation**

408 To estimate an oculomotor plant model for each larva, we used measured step responses to 409 calculate parameters of the linear filter given by equation (2) above. We assumed that each 410 step response was the result of an applied force convolved with a linear filter representing the oculomotor plant. As described above, a linear filter model implies that the eye position 411 412 after release from displacement will have the same number of components and the same time 413 constants as the plant (see equation (3)). However, the coefficients of the plant $\{c_i\}$ must still 414 be determined. Because the profile of the applied force is unknown, finding the coefficients 415 of the plant requires simultaneously finding the appropriate applied force profile. This is a 416 "blind deconvolution" problem and is generally under-constrained. Here, however, we have

417 two important constraints that facilitate finding a solution: first, we partially know the applied 418 force profile, i.e., that it is zero after the time of release; and second, the plant is assumed to 419 be a sum of exponentials with known time constants.

420 For each larva, we assembled eye position time series y_i starting from the onset of displacement for each response *i* until 8 to 14 s after release. All of the anaesthetized larvae 421 422 had 2 responses each, 3 of 6 awake larvae had two responses, and the remaining three had a 423 single response. Eye position was considered to be constant during displacement (see Step 424 response fitting). For awake larva time series, we used estimated step responses $y_{sr}(t)$ 425 resulting from both best and conservative extrapolations, as described above. Similarly to 426 fitting step responses, we removed the first 230 ms of data after release from displacement 427 due to ringing/oscillation shortly after release. Data points in this epoch were replaced with 428 the prediction of the best fit exponential model to the step response.

For linear filter estimation, we then defined the following loss function,

429

430
$$E(f_1, \dots, f_m, c_1, \dots, c_n) = \frac{1}{2} \sum_{j=1}^m \sum_{k=1}^{T_j} w_{jk} [(f_j * p(c_1, \dots, c_n))(t_k) - y_j(t_k)]^2.$$
(9)

431 Here, *m* is the number of responses for the larva, $f_j(t)$ is the time-varying force applied for 432 response *j*, $p(t; c_1, ..., c_n)$ represents the plant model parameterized by variable coefficients 433 { c_i } and fixed time constants { τ_i }, time is discretized as $t_k = k\Delta t$ where $1/\Delta t$ is the sampling 434 rate, and T_j is the number of time points in the response y_j . Each data point in the loss function 435 is weighted by a factor

436
$$w_{jk} = \begin{cases} 1/T_{\text{release},j} & \text{during displacement, } k \le T_{\text{release},j} \\ 1/(T_j - T_{\text{release},j}) & \text{after release} \end{cases}$$

437 where $T_{\text{release}, j}$ is the number of data points in the displacement period. This weighting causes 438 the loss function to be a sum of the mean squared fit error during displacement and the mean 439 squared fit error during the step response, with errors in these periods weighted equally even 440 though the period lengths are heterogeneous. We approximated convolution of continuous 441 signals with a discrete convolution, since Δt is small, bioRxiv preprint doi: https://doi.org/10.1101/2021.06.30.450653; this version posted July 1, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

442
$$(f * p)(t_k) = \Delta t \sum_{k'=0}^k f(t_{k'}) p(t_{k-k'}).$$
(10)

The blind deconvolution can then be computed by finding the forces $\{f_j\}$ and plant coefficients $\{c_i\}$ such that the loss *E* is minimized. First, we used the nonlinear solver TNC from 100 starting points that consisted of $\{c_{i,\text{initial}}\}$ chosen uniformly at random on (0, 1) and normalized to sum to 1. These preliminary results were used as the initial point, $\{c_i^{(0)}\}$ for an alternating least squares procedure, as follows:

448 1. Applied force estimation step. On iteration l + 1, given estimates for the plant 449 coefficients $\{c_i^{(l)}\}$, solve the optimization problem

450
$$f_1^{(l+1)'}, \dots, f_m^{(l+1)'} = \operatorname{argmin}_{f_1, \dots, f_m} E(f_1, \dots, f_m, \{c_i^{(l)}\})$$
subject to $f_i(t > t_{\text{release}}) = 0$ for $j \in \{1, \dots, m\}$

451 2. *Plant estimation step*. Given these estimates of the applied forces, update the estimate452 of the plant coefficients,

453
$$c_{1}^{(l+1)'}, \dots, c_{n}^{(l+1)'} = \operatorname{argmin}_{c_{1},\dots,c_{n}} E(\{f_{j}^{(l+1)'}\}, c_{1},\dots,c_{n})$$
subject to $c_{i} \ge 0$ for all *i*.

454 Note that the solutions obtained by this procedure are not unique: for example, if all 455 the plant coefficients are divided by a constant *K*, then as long as the resulting applied 456 forces are also multiplied by *K*, the new scaled coefficients still solve the problem. 457 We resolve this degeneracy by imposing the condition that the final plant coefficient 458 estimates sum to 1:

459
$$c_i^{(l+1)} = \frac{c_i^{(l+1)'}}{K} \text{ and } f_j^{(l+1)} = K f_j^{(l+1)'}, \text{ where } K = \sum_{i=1}^n c_i^{(l+1)'}.$$

These two steps are then repeated until the decrease in error is smaller than a threshold, $E^{(l+1)} - E^{(l)} \le 10^{-8}$. Note that the error cannot increase on any step, because in the worst case the same solution as the previous iteration can be chosen. Because the discrete convolution given by equation (10) is linear, we can write it as a matrix-vector product, and 464 each of the two steps above can be solved efficiently and accurately by a linear least squares
465 solver (lsq_linear in SciPy's "optimize" library). The plant model for each larva was chosen
466 to be the one of the 100 solutions that had the smallest final value for *E*.

We repeated this procedure using time constants and eye positions resulting from the conservative extrapolations in awake larvae. In order to evaluate the necessity of long time constants, we also repeated the procedure using only the fast (< 1 s) time constants resulting from the step response fit, both for anaesthetized and awake larvae.

For each plant model, we calculated the fractional contribution of each exponential component to the total area under the curve by integrating over the range t = 0 to t = 60 s for anaesthetized plants, and to t = 20 s for the active state plant models.

474

475 Neural drive estimation

476 Using the active state plant models, we estimated the force needed to produce the eve position 477 profile observed during fixation. We assumed this force was proportional to the neural drive 478 output by motor neurons. We computed neural drive estimates by deconvolving active state 479 eye position during the period of fixation preceding displacement with the corresponding plant model for each larva. In order to reduce noise, we used $y_{target}(t)$ as a smoothed 480 481 representation of eve position before displacement (see Fitting step responses), and 482 performed the deconvolution by solving a linear least squares problem with non-negative 483 least squares, as described above (Oculomotor plant model estimation). We measured the 484 persistence of both the estimated neural drive and the corresponding eye position by summing 485 the time series during fixation starting from and normalized to the value at 0.25 s post-486 saccade, and dividing this sum by the number of elements in the time series (Lee et al. 2015). 487 With this measure, a perfectly stable time series would have a persistence value of 1.

We also directly calculated the time constants of the slide components of the neural drive for each plant by calculating the poles of the Laplace transform of the neural drive. This was equivalent to finding the roots of the polynomial

491
$$N(s) = \sum_{i=1}^{n} c_i \prod_{j \neq i} (s + k_j),$$
(11)

19

492 where c_i and k_i are the amplitude and inverse time constant of the *i*th plant component, as in 493 equations (1) and (4) (see Mathematical Appendix, Analytical calculation of neural drive for 494 a multiexponential plant, for derivation). This was done numerically using the function 495 "roots" in NumPy.

496

497 Comparison of estimated neural drive to hVPNI activity

498 *Optical recordings* To compare the required neural drive to the activities of cells in the 499 hVPNI, we used previous optical recordings of somatic calcium-sensitive fluorescence in a 500 separate set of 6 larvae to estimate neuronal firing rates during fixations (Miri et al. 2011b). 501 Calcium-sensitive dye loading and optical recording methods are described in the original 502 reference. Data were collected using Oregon Green BAPTA-1 AM (Invitrogen) on a custom-503 built laser-scanning two-photon microscope that allowed synchronous eye tracking and 504 fluorescence image time series collection from sagittal planes within the hindbrain. 505 Fluorescence data acquisition and microscope control were performed using Cfnt v.1.529 506 (Michael Mueller, MPI, Heidelberg). Images were 256 x 256 pixels spanning 100 µm x 100 507 μ m regions and acquired at 2 ms per line (~2 Hz) in time series of 750 frames. For each larva, 508 five or six fluorescence image time series were collected from image windows lying in 509 parasagittal planes at fixed dorsoventral and rostrocaudal coordinates in the ventral $\sim 2/3$ of 510 the caudal hindbrain (rhombomere 7/8). All data analysed here were collected in the dark to 511 eliminate visual feedback.

512

513 hVPNI firing rate estimation For each cell, we determined baseline fluorescence to be the 514 smaller of two quantities: the mean of the saccade-triggered average fluorescence from 2 s 515 to 1 s before ipsiversive saccades, and the mean of the saccade-triggered average 516 fluorescence from 4 s to 5 s after contraversive saccades. We fit a calcium impulse response 517 function (CIRF), modelled by a single exponential decay, to baseline-subtracted 518 contraversive saccade-triggered average fluorescence time series, as in previous studies 519 (Daie et al. 2015; Miri et al. 2011b) and calculated the coefficient of determination, R^2 , for each fit. As in previous studies (Miri et al. 2011a, Miri et al. 2011b), cells for which CIRF 520 fits had $R^2 \ge 0.5$, and the Pearson correlation between saccade-triggered average fluorescence 521

and CIRF-convolved eye position after a contraversive saccade was > 0.5, were used (167 of
195 cells).

For each cell *j* satisfying these criteria, we modelled baseline-subtracted ipsiversive saccade-triggered average fluorescence x_j as a convolution between the CIRF and the sum of two components: a delta function modelling the saccadic burst and a multiexponential function representing the saccade-triggered average firing rate,

528
$$x_{j}(t) = e^{-k_{\text{CIRF},j}t} * \left(a_{0}\delta(t) + \sum_{i=1}^{n}a_{i}e^{-k_{i}t}\right) + \varepsilon_{j}(t), \qquad (12)$$

529 where $k_{\text{CIRF},i}$ is the cell-specific inverse CIRF time constant, k_i are inverse time constants of the saccade-triggered average firing rate, with coefficients $a_i \ge 0$ for all components *i*, and 530 ε_i is Gaussian noise with mean 0 and variance σ_i^2 . For each number of exponential 531 components *n* between 1 and 3, we solved for the coefficients $\{a_i\}$ and inverse time constants 532 533 $\{k_i\}$ that minimized the mean squared error of the model fit to data, using the nonlinear solver 534 TNC from 100 initial sets of parameters. For each number of components n, we picked the 535 best *n*-component model fit to be the one with the smallest mean squared error. Separately 536 for each cell, we called n^* the value of *n* after which adding another component decreased 537 the mean squared error by less than 1%. Then, we defined the best overall firing rate model 538 fit for each cell to be the best n^* -component model fit. We analysed responses from cells 539 where the ratio of the sum of squared errors of the best firing rate model fit to the sum of 540 squares of the data (sum of squares ratio, SSr) was less than 0.007 (151 of 167 cells)—cells 541 with SSr greater than this value had fits that were notably worse upon visual inspection. For 542 each cell, we calculated firing rate persistence values in the same manner described above 543 for neural drive and eye position persistence (Neural drive estimation).

544

545 *Summary plant and neural drive estimation* For the 9 active state step responses from 6 546 larvae, we simultaneously fit a single set of time constants, using the same procedure as above 547 (Step response fitting). Then, using these time constants, we performed the blind 548 deconvolution procedure (Oculomotor plant model estimation) to find the best fit plant model for all 9 responses simultaneously ("summary plant"). We then computed the saccadetriggered average eye position using smoothed versions of fixations from each of 6 separate larvae in which optical recording was performed. Smoothing was performed by fitting multiexponential functions to the saccade-triggered average eye position from each animal, as described above (Step response fitting). A neural drive estimate was then generated for each larva by deconvolving the saccade-triggered average eye position with the summary plant.

556 We then performed a regularized linear regression of the saccade-triggered average 557 firing rates of all hVPNI neurons onto the neural drive estimate for each larva, with regression 558 weights constrained to be nonnegative, using a linear least squares solver (lsq_linear). We 559 used a regularization term equal to the sum of squares (L_2 norm) of the regression weights of 560 each time series, multiplied by a regularization parameter. We chose this parameter 561 separately for each larva by bisection search so that approximately 50% of the weights were 562 nonzero, broadly in agreement with the fraction of putative hVPNI neurons that synapse onto 563 motor neurons in electron microscopy data (Lee et al. 2015, Vishwanathan et al. 2017).

564 To show the importance of intermediate timescales in the neural drive (Mathematical Appendix), for each larva we generated 100 synthetic populations of 150 mock cells whose 565 firing rates were each described by an exponential decay. For each population, 75 of the cells' 566 567 firing rates had time constants randomly drawn uniformly between 0 and 1 s, and 75 had time constants generated by taking 10 to the power of a uniform random number between 1.3 and 568 569 2, resulting in random time constants between 20 s and 100 s. We then performed a 570 regularized linear regression of the mock cells' firing rates onto the estimated neural drive 571 exactly as for the real cells.

572 **RESULTS**

573 Measurement of zebrafish oculomotor plant response under anaesthesia

574 All experiments described here were performed in the dark to prevent the influence of visual 575 feedback. In order to determine whether the larval zebrafish oculomotor plant, like that of 576 the anaesthetized primate, shows both short and long response timescales (Sklavos et al. 577 2006; Sklavos et al. 2005), we first measured the response of the eye to horizontal step 578 displacements (referred to below as "step responses") of two different durations in larvae 579 anaesthetized with MS-222. Because MS-222 inhibits action potential firing and thus input 580 to neuromuscular junctions, active muscle tone is likely diminished or absent in this state so 581 that the observed responses primarily reflect the passive properties of the plant. Similar 582 amplitude abducting step displacements lasting 10 and 60 s were applied to one eye of 583 anaesthetized larvae (n = 10) and the return trajectory of the eye was tracked following 584 release (Figure 2A,B). The use of two different step durations helps expose both short and 585 long response timescales; if the plant responded on exclusively short timescales, it would 586 effectively reach steady state within 10 s, so that the responses to 10 and 60 s displacements 587 would be similar.

588 However, the responses to 10 and 60 s step displacements were strikingly different 589 (Figure 3). To quantify response timescales present in these trajectories, for each larva we 590 simultaneously fit with multiexponential functions the first 60 s following release of 591 responses to both displacements. These functions had from 1 to 6 exponential decay 592 components and were constrained at the time of release to equal the eye position prior to 593 release. For each function with a given number of components, fits assumed a single set of 594 time constants but distinct sets of corresponding amplitudes for the responses to the 10 and 595 60 s displacements. Oscillatory artefacts sometimes present in responses during the first 200 596 ms following release, perhaps resulting from the manual control of the hydraulic manipulator, 597 led us to omit the first 230 ms of responses when fitting. Thus, our model fits cannot be 598 expected to well capture response timescales faster than ~ 100 ms. Single exponential 599 functions failed to capture much response structure, while models with 4 components 600 provided a better fit (Figure 3A,B). Improvements in fit quality as the number of model 601 components increased were similar to those observed previously for plant responses

measured in anaesthetized monkeys (Sklavos et al. 2005) and awake mice (Figure 3B; Stahlet al. 2015).

604 In order to determine the minimum number of discernible timescales in step 605 responses, we examined the change in mean squared error of fits as the number of exponential 606 decay components used in the fits was increased. For each larva, there was an "elbow" at 4 607 components, after which the reduction in mean squared error by adding more components 608 was extremely small (Figure 3B). The median values of the time constants of the four-609 component fits were 0.092, 1.34, 7.95 and 91.6 s (Figure 3C). Bootstrapped variance 610 estimates for time constant values were small relative to the gaps between values for 611 successive components (median sd of estimates ranged between 2.1 and 6.6% as a fraction 612 of the time constant values). Time constants greater than 10 s were necessary to well fit 613 responses; multiexponential models constrained to have time constants less than 10 s failed 614 to fit step responses as well as unconstrained models having equivalent numbers of 615 components (Figure 3A; 39- to 172-fold increase in MSE for four-component models).

616 We next measured step responses under another anaesthetic, the NMDA receptor 617 antagonist ketamine. These experiments were performed for three reasons: (a) to demonstrate 618 that aspects of the results obtained with MS-222 are not dependent on the choice of 619 anaesthetic, (b) to better compare our results in the larval zebrafish with those of Sklavos et 620 al. (2005) in the primate, and (c) because there is some indication in mammals that at 621 ketamine doses near the threshold above which the animal becomes unresponsive to eye 622 manipulation, some active muscle tone is preserved, making ketamine anaesthesia potentially 623 closer to the active state (Blanks et al. 1977; King et al. 1978; Sklavos et al. 2005). We did 624 not attempt to verify the presence of active muscle tone under ketamine. After anaesthetizing 625 larvae (n = 6) with ketamine, we applied similarly sized abducting step displacements of 15 626 and 90 s to one eye and tracked its return trajectory following release (Figure 3D).

We simultaneously fit the responses following 15 and 90 s displacements using multiexponential functions as described above. The choice of slightly different step durations here was arbitrary and should not obscure the general agreement this similarity demonstrates. Fit results were similar to those obtained under MS-222 (Figure 3D-F). The median values of the time constants of four-component fits were 0.131, 2.01, 10.7, and 110.2 s, similar to those obtained under MS-222 (Figure 3F). Bootstrap confidence interval estimates for time constant values were small relative to the gaps between values for successive components (median sd of estimates ranged between 4.7 and 8.4% as a fraction of the time constant values). Models constrained to have time constants less than 10 s again failed to fit step responses well (Figure 3D, 8- to 58-fold increase in MSE for four-component models).

637 Collectively, the data from anaesthetized larvae suggest that the larval zebrafish 638 oculomotor plant, like that of the primate, demonstrates both short (< 1 s) and long (> 1 s) 639 response timescales. Moreover, our observation of time constants spread over several orders 640 of magnitude under both types of anaesthesia matches results from comparable primate 641 experiments (Sklavos et al. 2006; Sklavos et al. 2005).

642

643 Measurement of oculomotor plant responses in the active state

644 Because active tone in extraocular muscles could influence the response properties of the 645 oculomotor plant, we examined whether long response timescales were discernible in awake, 646 behaving larvae. Previous observations of active state plant responses have been limited to 647 relatively brief time windows (< 400 ms in monkey, Anderson et al. 2009; < 2 s in mouse, 648 Stahl et al. 2015), potentially obscuring the presence of long response timescales. Here we 649 succeeded in measuring active state plant responses of longer duration thanks to the relatively 650 low saccade frequency of larval zebrafish. Five to seven seconds after an adducting saccade 651 in the dark, we applied abducting step displacements lasting 6.5 to 8.5 s. On 9 occasions 652 across 6 larvae, we were able to record responses lasting > 8 s without any interrupting 653 saccades. To our knowledge, active state measurements of comparable duration have not 654 been previously reported.

In order to properly fit these responses, we needed to estimate what the eye position would have been during the responses had the imposed displacements not occurred (Figure 2C). Because eye position decays toward the centre of gaze appreciably during fixations in larval zebrafish, we could not use the eye position immediately prior to displacement as an estimate of the expected eye position in the absence of displacement. Instead, we estimated the expected eye position by fitting multiexponential models to eye position between the previous adducting saccade and the displacement, and then extrapolated the model fits 662 forward in time to estimate eye position in the absence of displacement. Models having 663 between 1 and 6 components were initially fit, and the number of components used for 664 subsequent analysis was chosen using the reduction in mean squared error from each added component as described above. In addition to this "best" extrapolated eye position fit, we 665 also performed a second "conservative" fit with faster eye position decay (see Methods, 666 667 Figure 2D) that conservatively accounts for possible over-estimates in the duration of the 668 step response resulting from extrapolation. We extrapolated both the "best" and 669 "conservative" fit functions through the step and subsequent response epochs to predict eye 670 position had the displacement not been applied. We defined the step response as the 671 difference between the eye position following release and these extrapolated fit functions 672 (Figure 2C, grey arrows).

673 To quantify response timescales in the active state, we fit multiexponential functions 674 to responses from 230 ms to 8 - 21 s following release for each larva. For 3 larvae, a single 675 response was fit. For 3 other larvae, responses to two separate displacements were 676 simultaneously fit with a common set of time constants, but distinct component amplitudes 677 for each displacement were permitted. Results assuming the best and conservative 678 extrapolations were generally in agreement. Single exponential functions failed to well-679 capture response structure (Figure 4A,B). For 5 of 6 larvae, responses were best fit by a three-680 component model (Figure 4C), while responses from the remaining larva was best fit by a 681 four-component model. Fit improvements upon inclusion of additional components beyond 682 4 were again generally very small (Figure 4D). Time constants were broadly distributed for 683 all larvae, assuming both the best and conservative eye position extrapolations (Figure 4E,F). 684 Assuming the best extrapolation, the median values of the time constants of three-component 685 fits for all 6 larvae were 0.166, 2.17, and 36.3 s. Bootstrapped variance estimates for time 686 constant values were again small relative to the gaps between values for successive 687 components (median sd of estimates was around 5% as a fraction of the inverse time constant 688 values). Here again, very long time constants were necessary for an adequate fit; multiexponential models constrained to have time constants no greater than 5 s produced 689 690 much worse fits to step responses than unconstrained models having equivalent numbers of 691 components (Figure 4A,B, 3- to 459-fold increase in MSE for three-component models using the best extrapolation and 1.3- to 356-fold increase using the conservative extrapolation).
Collectively, these results demonstrate that, even when accounting for possible errors in
extrapolation, active state step responses also display both short and long response
timescales.

696 Comparing the active state results to those obtained with anaesthesia reveals strong 697 similarities. The mean time constants across larvae from active state fits are qualitatively 698 similar to those found under both MS-222 and ketamine (Figure 3C,F), again ranging over 699 several orders of magnitude. Quantitatively, the time constants for the active state fits tended 700 to be a few times smaller than for the anaesthetized preparations. However, this may be an 701 artefact of the limited measurement time in the active state; the mean time constants we found 702 when we fit only the first 15 s of post-release eye position in anaesthetized larvae differed 703 from those of the best 3 component fits to active state larvae by just 10-20%.

704

705 Implications for neural drive

706 We next addressed the implications of our above plant response observations for the nature 707 of the neural drive extraocular muscles would require in order to stabilize gaze during 708 fixation. To estimate this neural drive from eye position measurements, we used a simplified 709 model that captures salient plant response properties. Our model assumes that eye position 710 can be approximated by a convolution of neural drive with the plant's response dynamics. 711 We used the active state step responses to calculate the parameters of a linear filter 712 representing the impulse response function of the oculomotor plant for each larva. To 713 compute these plant models, we calculated filters that would best map a force applied to the 714 eye during displacement onto the response observed after release (see Methods, Figure 5A). 715 The response after release for a multiexponential plant model is a multiexponential decay 716 with time constants equal to those of the plant model. Thus, we defined "distributed" plant 717 models as multiexponential filters constrained to have the same number of components and 718 time constant values as the fits to active state step responses.

719 We observed three pertinent features of the resulting plant models. First, we found 720 that models derived from both the best and conservative extrapolations of eye position could 721 accurately reproduce the active state step responses (best extrapolation: mean $R^2 \pm sd = 0.965$ ± 0.032 , conservative extrapolation: mean R² \pm sd = 0.948 \pm 0.054; Figure 5B-D). Second, the functions inferred for force during displacement appeared to be composed of pulse, step, and some number of exponential slide components, as expected under a multiexponential plant model (Figure 5C, red; Mathematical Appendix). Lastly, plant model amplitudes were largest for the fastest components (Figure 5D,E). With component amplitudes summing to one, the median amplitude for the fastest component was 0.922 or 0.917 assuming the best or conservative extrapolations, respectively.

729 Given the small relative amplitudes of the long timescale components, we asked what 730 impact these components would make on gaze stabilization during fixation. Despite their 731 small amplitudes, these components contribute a comparatively large fraction of the area 732 under the curve of the plant model response functions up to 20 s post-impulse (Figure 5F), 733 suggesting an important role in governing plant responses. To verify this, we repeated the 734 plant model estimation using multiexponential filters in which the components with time 735 constants greater than 1 s were removed, leaving the one or two fastest components ("fast 736 plant" models). The remaining time constants align well with those of previous models of 737 the mammalian oculomotor plant (Goldstein 1984; Optican and Miles 1985; Robinson 1964; 738 Sklavos et al. 2006; Stahl and Simpson 1995; Stahl et al. 2015). However, using filters 739 lacking long timescale components resulted in much poorer step response estimates (best extrapolation: mean $R^2 \pm sd = -0.136 \pm 0.068$, conservative extrapolation: mean $R^2 \pm sd = -$ 740 0.191 ± 0.090 ; Figure 5B). Results were similar using step response fits where time constants 741 742 were constrained to be less than 1 s. Thus, long timescale components with even the small 743 relative amplitudes we observed appear to play a critical role in dictating plant responses.

744 The above observations rely on our measurements of active state step responses, 745 which themselves involved the extrapolation of eye position expected in the absence of 746 displacement. Therefore, we checked to see if substantially similar plant models are 747 consistent with step responses measured under anaesthesia, which involved no extrapolation 748 of eye position (Figure 6A). We again found that our models could reproduce measured step 749 responses (Figure 6B,C), that inferred force functions were composed of pulse, step, and 750 exponential slide components (data not shown), and that the fastest components had the 751 largest amplitudes (Figure 6D). Importantly, the distributions of component time constants,

amplitudes, and fractional area under the curve for the inferred plant models (Figure 6D,E)
were similar to the distributions observed for active state plant models (Figure 5E,F). Thus,
the pertinent features of recovered plant models do not appear to be artefacts of our use of
extrapolation in active state response measurements.

756 Based on the estimated active state plant models, we next estimated the neural drive 757 required to stabilize gaze during fixation (Figure 7A-D). We focus here on the neural drive 758 needed long after the saccade has ended (> 0.25 s after initiation) to distinguish this drive 759 from the large transient drive needed to quickly move the eye during a saccade. In the case 760 of fast plant models that lack long timescale components, the neural drive needed to fully 761 stabilize eye position is essentially constant over this epoch (Figure 7B, blue dashed line). 762 This reflects the step component (Goldstein and Robinson 1986; Optican and Miles 1985; 763 Robinson 1964) that has long been thought to be necessary for gaze stability during fixation, 764 representing the temporal integral of eye velocity commands. The time course of this estimate 765 is not dependent on the precise values of filter parameters – essentially constant drive will be 766 necessary to stabilize eye position whenever filter time constants are exclusively fast 767 (Mathematical Appendix).

768 The addition of multiple long timescale components to the plant model markedly 769 changes the character of the neural drive required to fully stabilize gaze (Figure 7B, red 770 trace). First, the drive is no longer dominated by a step component, as substantial decay in 771 drive provided by slide components is seen seconds into the fixation. This makes intuitive 772 sense, as the step component is filtered by the plant's long timescale components, so the step 773 amplitude required is smaller. In fact, the step component amplitude can be shown to be 774 inversely proportional to the area under the plant filter (Mathematical Appendix). Second, 775 slow decay in the neural drive, generated by the slide components, is necessary to compensate 776 for the slow equilibration of the viscoelastic elements to their steady states. We directly 777 calculated the slide component time constants, finding that exactly one fell between each pair 778 of consecutive plant model time constants and at least one slide time constant was always > 779 1 s (Figure 7D). To test whether slide time constants fall between consecutive plant model 780 time constants more generally, we directly calculated slide time constants for simulated three-781 or four-component plants and found that the observation held over the broad range of parameters tested (Mathematical Appendix). Thus, the presence of long timescale plant
responses appears to imply the presence of slowly decaying neural drive components.

784

784 Actual measurements of larval zebrafish eye position show prolonged but imperfect 785 fixations (Figure 7C, grey trace). Nevertheless, neural drive estimates obtained using 786 distributed plant models, unlike those obtained using fast plant models, again decay more 787 quickly than eye position (Figure 7C, red vs. blue trace). Furthermore, the neural drive 788 contains long timescale slide components identical to those required to maintain perfect 789 fixation (Figure 7D). We show in the Mathematical Appendix that, under the inverse model 790 formulation, neural drive yielding imperfect fixations more generally has slide components 791 identical to those that generate stable fixations, in addition to components resembling 792 recorded eye position that are analogous to the step component for stable fixations.

793 How much do neural drive estimates for distributed plant models differ from those 794 for fast plant models, given actual eye position measurements? We used "persistence values" 795 to quantify the difference in time course persistence expected between neural drive and eye 796 position, assuming either distributed or fast plant models (9 fixations from 6 larvae; Lee et 797 al. 2015). We defined the persistence value for a neural drive or eye position time series as 798 its integral starting from 0.25 s post-saccade, normalized so that a stable time series yields a 799 persistence value of 1. Hence larger persistence values correspond to increasingly persistent 800 time courses. Distributed plant models lead to neural drive persistence that is substantially 801 less than eye persistence, whereas fast plant models imply similar drive and eye persistence 802 (Figure 7E). The median ratios between the persistence values of estimated neural drive and eye position were 0.72 and 1.04 for the distributed and fast plant models, respectively. Thus, 803 804 distributed plant models predict neural drive will substantially diverge from eye position.

805

806 The relationship between measured hVPNI activity and eye position

We next assessed whether the activity of neurons in the hVPNI, which has traditionally been assumed to generate a constant step component, is consistent with a role in stabilizing gaze given the long response timescales in the plant and their implications for neural drive outlined above. We first compared the difference in time course persistence expected between the neural drive and eye position during fixation for individual larvae (Figure 7E) with that seen 812 between measurements of hVPNI firing and eye position recorded in a previous study (Miri 813 et al., 2011a,b). Time course persistence was computed for hVPNI neurons whose saccade-814 triggered average firing rates were estimated from calcium-sensitive cellular fluorescence 815 measured during saccadic eye movement. We modelled saccade-triggered average 816 fluorescence as a multiexponential firing rate function convolved with a calcium impulse 817 response function describing the fluorescence response following an action potential (see 818 Methods, Figure 8A,B; Miri et al. 2011a,b; Daie et al. 2015). Out of an initial dataset of 195 819 neurons across 6 larvae, we excluded neurons that were not putative integrator neurons 820 (28/195 neurons excluded) such as those exhibiting bursting activity in response to a saccade, 821 and those for which fluorescence was not well fit by the saccade-triggered average 822 fluorescence model (16/195 neurons excluded), leaving 151 neurons for subsequent analysis. 823 Consistent with previous results, we observed a broad range of persistence values across this 824 population (Figure 8C).

825 We then compared persistence values for hVPNI neurons with those of the saccade-826 triggered average eye position calculated from simultaneously recorded eye position (Figure 827 8D). We found that firing rate and eye persistence measurements were statistically unlikely 828 to be drawn from the same distribution (p = 0.0106, two-sided Wilcoxon rank-sum). Firing 829 rate persistence was lower than that for the corresponding eye position for 122/151 neurons 830 (81%). The median ratio between the persistence values for firing rate and eye position across 831 all neurons was 0.69, very close to the corresponding value for neural drive estimates (0.71, 832 red line in Figure 8D). Overall, the distribution shown in Figure 8D is more consistent with 833 what would be expected given the distributed plant models than given the fast plant models.

834 We next addressed whether the firing rates of the population of hVPNI neurons could 835 be used to construct a neural drive that stabilizes gaze given long response timescales in the 836 plant. For this analysis, we generated a three-component summary plant model computed as 837 for the individual larvae but using a simultaneous fit to the active state responses from all six 838 larvae (Figure 9A-C). This summary model had component amplitudes and time constants 839 similar to the medians of the corresponding distributions from models for individual larvae 840 (Figure 9B). We then deconvolved the saccade-triggered average eye position for each larva 841 with this summary plant model to generate a neural drive estimate. Assuming that hVPNI 842 output could effectively be fed forward by motor neurons, we asked whether weighted sums 843 of saccade-triggered average firing rates for recorded hVPNI neurons could well-844 approximate these neural drive estimates (see Methods, Figure 9D-F). Weights were constrained so that 50% were positive, and the rest zero, in agreement with anatomical 845 846 estimates of the proportion of integrator cells that synapse onto motor neurons (Lee et al. 847 2015, Vishwanathan et al. 2017). We found that a weighted sum of hVPNI firing could wellapproximate the neural drive estimate for all 6 larvae ($R^2 = 0.989-1.000$). This indicates that 848 hVPNI firing is sufficient to constitute the neural drive needed to stabilize an oculomotor 849 850 plant characterized by long response timescales.

That such good fits can be achieved is perhaps not surprising given the range of persistence timescales present in hVPNI activity (Figure 8C). However, we note that intermediate timescales are critical to fit neural drive estimates well. Weighted sums of simulated cell populations containing only a distribution of short (<1 s) and very long (>20 s) timescales could not reconstruct neural drive estimates (Figure 9E, cyan traces; 10- to 183fold increase in MSE). Thus, the intermediate persistence timescales not present in classical models of the oculomotor neural integrator are a necessary component of this drive.

858 **DISCUSSION**

859 We report here two significant findings regarding the oculomotor plant and the motor circuits 860 that control it. First, we extend the demonstration of both short and long response timescales 861 in the plant (Quaia et al. 2009; Sklavos et al. 2006; Sklavos et al. 2005) to the active, 862 unanaesthetized state, and to a new vertebrate model organism. Recent reports of long 863 timescale responses in the primate plant have been conducted under anaesthesia. The 864 relevance of these responses to the active state has been argued only indirectly using models 865 (Sklavos et al. 2005). Second, our results establish that, despite such long timescales, the 866 firing seen among hVPNI neurons (Daie et al. 2015; Miri et al. 2011a) can still be interpreted 867 in terms of an inverse model-based compensation of plant viscoelasticity. While previous 868 work has focused on the encoding of eye position in neuronal populations necessary for gaze 869 stability during fixation (Aksay et al. 2000; Escudero et al. 1992; McFarland and Fuchs 1992; 870 Pastor et al. 1994), a deviation from simply representing eye position appears crucial to the 871 hVPNI's function. As predicted for an inverse plant model that achieves substantially stable 872 gaze, firing among larval zebrafish hVPNI neurons shows both less persistence on average 873 than eye position itself, and a heterogeneity of persistence timescales.

874 Previous models of the oculomotor plant that included only fast response timescales 875 implied that during fixation, neural drive to the plant would decrease over the first tens or 876 hundreds of milliseconds and thereafter would stably approximate eye position (Goldstein 877 1984; Optican and Miles 1985; Robinson 1964). This decrease in drive that follows the 878 saccade-inducing burst, attributed to an exponentially decaying slide component, reflects the 879 attenuating force needed to stabilize gaze as the viscoelastic elements equilibrate. The 880 presence of distributed response timescales in the plant that range up to tens of seconds 881 implies a reduced need for the constant, step component of neural drive; instead, distributed 882 timescales of decaying drive extend long into fixations to compensate for the distributed 883 timescales of force dissipation. Indeed, measurements of abducens motor neuron firing 884 during approximately stable fixations in cats show evidence of firing rate decay on timescales 885 greater than 1 s (Davis-Lopez de Carrizosa et al. 2011). Hysteresis observed between 886 abducens motor neuron firing and eye position greater than 2.5 s into fixations (Goldstein 887 and Robinson 1986) is also consistent with the presence of neural drive components on the many seconds timescale, as is the hysteresis seen across timescales between hVPNI neuron
firing and eye position (Aksay et al. 2003).

890 The requirement for neural drive that decays across short and long timescales complicates descriptions of drive to the plant. The notion that the drive is composed of an 891 892 eye velocity-encoding component, an eye-position component, and an exponentially 893 decaying slide component can be extended to include multiple decaying slide components 894 distributed across a range of timescales (see Mathematical Appendix). However, long 895 response timescales in the plant imply that the drive required for stable gaze will have a step 896 component with small relative amplitude. Furthermore, because sums of exponential 897 components differing in component number, time constants, and amplitudes can well-898 approximate the same function (Istratov and Vyvenko 1999), the presence of at least several 899 response timescales implies that discerning precise values for these parameters may not be 900 possible. However, parameters from the multiexponential fits we used to characterize plant 901 responses do meaningfully reflect the distribution of response timescales.

902 Despite this ambiguity regarding the neural drive needed to stabilize gaze, the need 903 for multiple slide timescales does entail a coherent view of the transformation performed by 904 the hVPNI in stabilizing gaze. Each slide timescale can be computed as a "leaky" integral of 905 a brief eye velocity-encoding burst – an imperfect integral that "leaks" away over time with 906 a particular time constant. The hVPNI can then be viewed as computing a sum of multiple 907 leaky integrals of eye velocity (Figure 1A). Individual hVPNI neurons may reflect distinct 908 combinations of these integrals in their firing, as would the ocular motor neurons they target. 909 Since leaky integration can also be expressed as a convolution with an exponentially 910 decaying filter, the hVPNI can be seen as convolving eye velocity with a multiexponential 911 filter. One practical manifestation of this leaky integration of eye velocity is that hVPNI firing 912 will decay faster than eye position (the pure integral of eye velocity), consistent with our 913 observations of hVPNI firing in the aggregate. While pure integration remains a useful 914 approximation of the transformation performed by the hVPNI, the distributed nature of its 915 integration timescales may have important consequences for the underlying biological 916 mechanisms (Daie et al. 2015; Miri et al. 2011a; Seung 1996; Seung et al. 2000). In previous

work, we have identified an array of neural circuit architectures capable of generating leakyintegrals on multiple, distributed timescales (Miri et al. 2011a).

919 This view of the hVPNI as decaying on multiple timescales is conceptually similar to 920 previous work suggesting power law decay of neural firing rates in the oculomotor integrator, 921 since power law decays can be approximated by weighted sums of exponential decay terms. 922 Previous work has proposed that the transformation of eye velocity signals by the hVPNI is 923 not pure temporal integration, but fractional-order integration (Anastasio 1994). What is 924 commonly referred to in calculus as integration (integration "of order 1") can be generalized 925 to integrations of arbitrary real (hence "fractional") order (Podlubny 1999). Integration of an 926 order between 0 and 1 is equivalent to convolution with a power function filter of the form 927 t^{α} , where α is equal to the integration order minus one. Finding evidence for integration of 928 order between 0 and 1 in oculomotor circuits, Anastasio (1994) posited that the neural drive 929 to the oculomotor plant may constitute a fractional integral of eye velocity. He further 930 proposed that this fractional integral relation might serve to compensate fractional integration 931 of neural drive by the plant. That is, the motor circuitry and the plant itself may each be 932 contributing a fraction of the integration necessary to transform eye velocity signals into a 933 position signal.

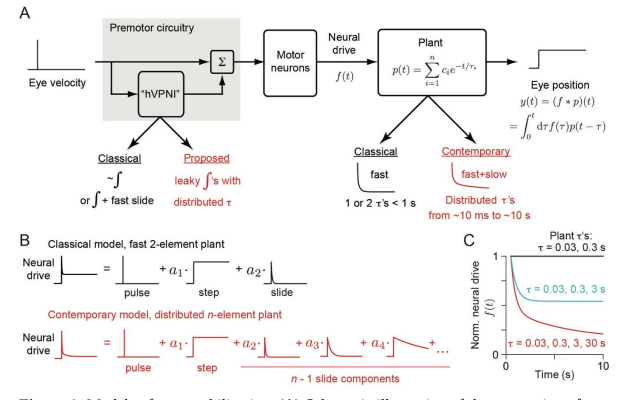
934 Our results here show that the range of persistence timescales present in hVPNI firing 935 is broad enough to constitute a signal matching estimates of the neural drive needed to dictate 936 eye position during fixation. Although contemporary multiexponential plant models imply 937 that this drive should differ from previous descriptions of hVPNI output emphasizing the 938 encoding of eye position, we show here that linear sums of larval zebrafish hVPNI neuron 939 firing, relayed by motor neurons, could stabilize gaze assuming the plant models we 940 computed. Observations of distributed persistence timescales in the hVPNI of adult goldfish 941 (Major et al. 2004; Miri et al. 2011a) and cat (Davis-Lopez de Carrizosa et al. 2011) suggest 942 that hVPNI output and oculomotor plant dynamics could be similarly reconciled for adult 943 vertebrates.

Despite the sufficiency of hVPNI firing for comprising a drive that can compensate for distributed response timescales in the oculomotor plant, the variation in persistence across neurons may still seem curious. If the plant is well modelled by a single filter, then there will 947 exist a single fixed mapping between eye velocity-encoding commands and the drive needed 948 to produce observed eye position, reflecting a single inverse model of the plant. This single 949 inverse model could be instantiated by the hVPNI such that all hVPNI neurons fired 950 identically, counter to our observations. One possible explanation is that the variation in 951 hVPNI persistence reflects the circuit architecture that gives rise to distributed persistence timescales, or an architecture that additionally enables context-dependence in the distribution 952 953 of these timescales across neurons, as has recently been observed (Daie et al. 2015). Another 954 possibility is that variation in neuronal persistence timescales provides a means for conferring 955 robustness to gaze control. Changes in plant dynamics could be counterbalanced simply by 956 reweighting inputs to motor neurons or the extraocular muscles, effecting a reweighting of 957 different timescales in the neural drive. Lastly, it is also possible that our measurements belie 958 a greater complexity in the plant. It could be that the plant comprises separate dynamic 959 components that are independently compensated by drive to extraocular muscles (Dietrich et 960 al. 2017; Hernandez et al. 2019). This would imply a need for premotor circuitry to instantiate 961 not one but multiple inverse models, and to generate distinct drive components that would not always be in similar proportion. Variation in the firing persistence of hVPNI neurons 962 963 would then be inevitable.

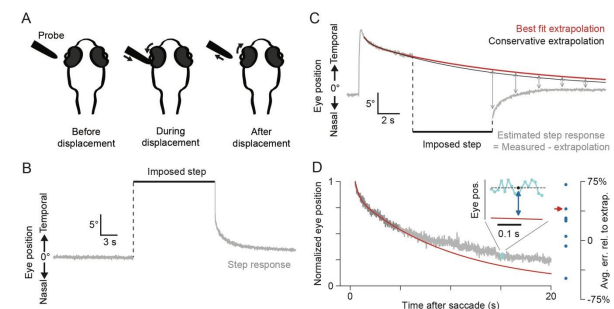
964 Our results reported here indicate that long response timescales are not unique to the 965 primate oculomotor plant. This adds to the demonstration by Davis-Lopez de Carrizosa et al. 966 (2011) that abducens motor neuron firing decays faster than abducens muscle tension during 967 fixation in cats, which suggests the presence of long response timescales in the plant as well. 968 Thus, the implications of these timescales for compensation by motor circuits may be further 969 addressed in a range of model organisms including the larval zebrafish, which is particularly 970 amenable to circuit-level analysis (Ahrens et al. 2012; Arrenberg and Driever 2013; Friedrich 971 et al. 2013; Orger et al. 2008) and has achieved prominence in the study of visuomotor 972 behaviour (Bianco et al. 2011; Gahtan et al. 2005; Helmbrecht et al. 2018; Okamoto et al. 973 2008; Portugues and Engert 2009; Sylvester et al. 2017). For example, how the cerebellum 974 contributes to oculomotor integration by tuning brainstem circuits could also be fruitfully 975 addressed using the larval zebrafish model (Miki et al. 2020; Aizenberg and Schuman 2011). 976 While gaze stability appears to require something beyond pure temporal integration, 977 the underlying circuitry remains a valuable model for circuit-level short-term memory (Major 978 and Tank 2004). In particular, the multiple timescales of firing persistence seen in the hVPNI 979 appear to be analogous to those seen in cortical short-term memory circuits (Bernacchia et 980 al. 2011; Machens et al. 2010), suggesting that multi-timescale responses may represent a 981 core feature of short-term memory systems throughout the brain. Moreover, models of neural 982 integration proposed in previous work on the larval zebrafish hVPNI (Miri et al. 2011a) have 983 been extended to interpret response diversity among oculomotor integrator neurons in 984 monkeys (Joshua et al. 2013). We note that leaky integration of velocity signals on distributed 985 timescales in the hVPNI still requires the generation of firing that persists much longer than 986 typical membrane and synaptic time constants. Cellular and/or circuit mechanisms must exist 987 that generate this persistence. Thus, elucidating these mechanisms in the larval zebrafish 988 should contribute to our understanding of short-term memory in other circuits.

989 FIGURES

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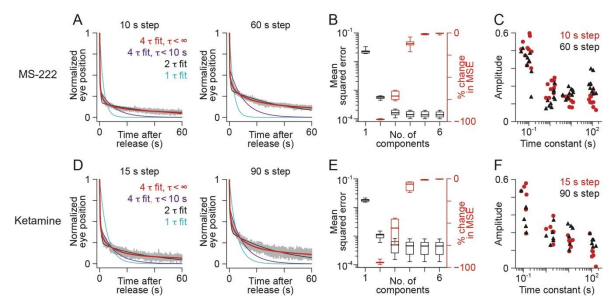
991 Figure 1 Models of gaze stabilization. (A) Schematic illustration of the conversion of eye 992 velocity commands into the neural drive f(t) necessary to maintain stable fixation. This neural drive is conveyed by the ocular motor neurons to determine eye position y(t). The 993 994 oculomotor plant is modelled as a linear filter p(t) operating on this drive. Premotor circuitry 995 generates components of f(t) from signals encoding eve velocity. The classical view, which 996 predates recent results, held that the plant can be characterized by components that relaxed 997 on 10's and 100's of ms timescales. Gaze stability on longer timescales then would require 998 the generation of a neural drive that approximates the temporal integral of eye velocity. Here 999 we argue that evidence for a broad distribution of response timescales in the plant redefines 1000 the role of premotor circuitry as involving a summation of leaky integrations on distributed 1001 timescales. hVPNI: horizontal velocity-to-position neural integrator. (B) Decomposition of 1002 the neural drive needed to stabilize gaze under classical and contemporary models of the 1003 oculomotor plant. (C) Neural drive required to maintain stable fixation, from 500 ms to 10 s 1004 after saccade termination, for a classical plant model (black) and ones with longer response 1005 timescales (cyan, red).



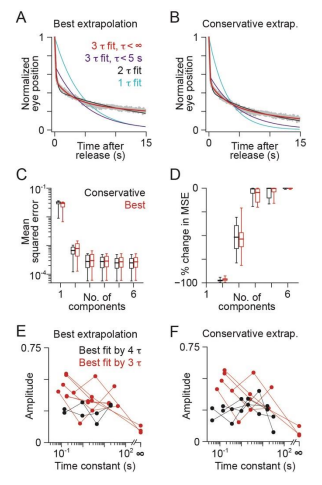


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1008 Figure 2 Methodological approach. (A) Schematic illustration of the method used to 1009 measure the oculomotor plant step response. A blunt probe controlled by a hydraulic 1010 micromanipulator was used to transiently displace the eye. (B) In anaesthetized larvae, the 1011 step response was the eye position measured after release from an imposed step displacement. 1012 (C) In awake larvae, the eve after displacement was assumed to return to the position it would 1013 have occupied had no displacement occurred. Thus, the step response was estimated to be 1014 the measured eye position after release from the step displacement (grey), minus the 1015 extrapolation of pre-displacement eye position (red). A faster decaying conservative 1016 extrapolation (black) was also calculated. (D) Left: to validate the extrapolations, fits to the 1017 initial portion (darker grey) of long unperturbed fixations were extrapolated (red), and the 1018 fractional error of the extrapolation relative to the true eye position was averaged over a ~230 1019 ms window centred at 14.64 s (inset: cyan, data; dashed black, mean of these data). Right: 1020 the average relative error across the 230 ms window for each measured fixation (red arrow 1021 corresponds to the fixation shown on the left).

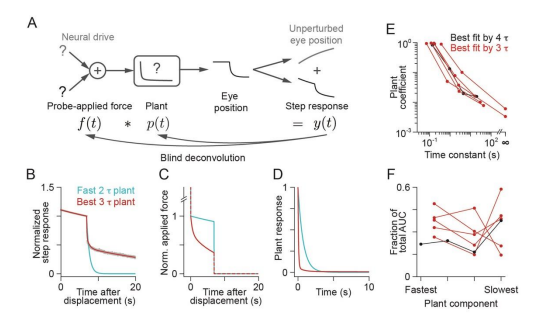


1024 Figure 3 The eye's return from step displacements in anaesthetized zebrafish larvae reveals 1025 short and long response timescales in the oculomotor plant. (A) Example responses of the 1026 eye (grey) following step displacements lasting 10 s (left) or 60 s (right), normalized by eye 1027 position prior to release, in larvae anaesthetized with MS-222. Coloured traces: simultaneous 1028 fits to both displacements with multiexponential models having one (cyan), two (black), or 1029 four components with time constants unconstrained (red) or constrained to be less than 10 s 1030 (purple). (B) Mean squared error (MSE) of fits with different numbers of components (black) 1031 and percent change in MSE compared to the fit with one fewer component (red) in larvae 1032 anaesthetized with MS-222. Boxes span the 25th to 75th percentile range; whiskers show 1033 maximum and minimum values (n = 10 larvae). (C) Coefficient amplitudes and time 1034 constants for four-component fits in larvae anaesthetized with MS-222. (D-F) Same as A-C 1035 but for larvae anaesthetized with ketamine (n = 6), for which step displacements lasted 15 1036 or 90 s.



1038 Figure 4 The eye's return from step displacements in awake zebrafish larvae exhibits short 1039 and long response timescales. (A,B) Example active state step response calculated using the best (A) and conservative (B) extrapolation of eye position in the absence of displacement. 1040 1041 Fits of multiexponential models having one (cyan), two (black) or three components with time constants unconstrained (red) or constrained to be less than 5 s (purple) are overlaid. 1042 1043 (C,D) Mean squared error (MSE) (C) and percent change in MSE (D) for step response fits 1044 compared to models with one fewer component, calculated using the best (red) or 1045 conservative (black) eye position extrapolations for each larva. Boxes span the 25th to 75th 1046 percentile range; whiskers show maximum and minimum values (n = 6 larvae). (E,F) Distributions of amplitudes and time constants for fits to step responses calculated using the 1047 1048 best (E) and conservative (F) eye position extrapolations. Dots and lines show parameter 1049 combinations for larvae for which the best fit was a three-component model (red; best

- 1050 extrapolation: n = 5, conservative extrapolation: n = 4) or a four-component model (black;
- 1051 best: n = 1, conservative: n = 2).



1052

1053 Figure 5 Oculomotor plant models can be recovered from step responses in awake zebrafish 1054 larvae. (A) Schematic outlining the method used to estimate linear filters that capture plant 1055 response properties. The measured eye position is assumed to derive from the sum of two 1056 inputs to the plant: external force applied by the probe, which leads to the active state step 1057 response y(t), and internally generated neural drive, which would lead to the (extrapolated) 1058 unperturbed eye position. We estimated the plant filter and applied force that best fit the 1059 active state step responses. Active state step responses were estimated from the "best" 1060 extrapolation procedure of Figure 2C and normalized to be 1 at the time of release. (B) 1061 Example active state step response (grey), and predicted step responses from the best recovered three-component plant model (red) or a plant model refit using only the fastest two 1062 1063 time constants from the three-component plant model (cyan). (C) Time course of recovered 1064 applied force, normalized to be 1 immediately after pulse offset, assuming the best three-1065 component (red) or fast two-component (cyan) plant. (D) Time course of the impulse 1066 response of the best three-component (red) and fast two-component (cyan) plant models. (E) 1067 Distribution of amplitudes and time constants for each component of the best recovered plant 1068 models. Dots and lines show parameter combinations for larvae for which a three-component 1069 model was best (red, n = 5) or for which a four-component model was best (black, n = 1). 1070 (F) Fraction of the total area under the curve (AUC), calculated over 20 s, contributed by

- 1071 each plant component for the best recovered three- (red, n = 5) or four-component (black,
- 1072 n = 1) plant models.

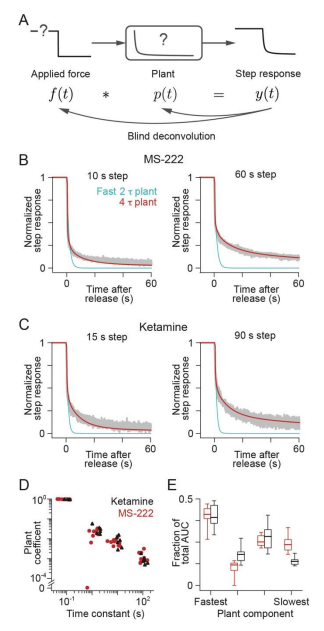
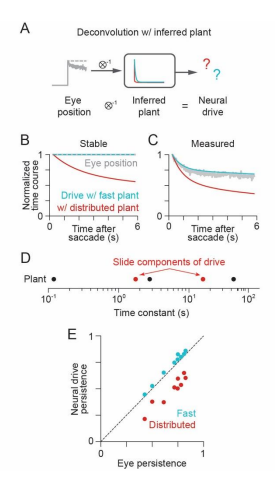


Figure 6 Oculomotor plant model estimates for anaesthetized larvae are consistent with active state estimates. (A) Schematic outlining the method used to estimate linear filters that describe plant response properties. We estimated the plant filter and applied force that best fit the measured anaesthetized step responses. (B) Example responses of the eye (grey) following step displacements lasting 10 s (left) or 60 s (right), normalized by the eye position prior to release, in larvae anaesthetized with MS-222. Coloured traces: predicted step response from the best recovered four-component model (red) and a plant model refit using

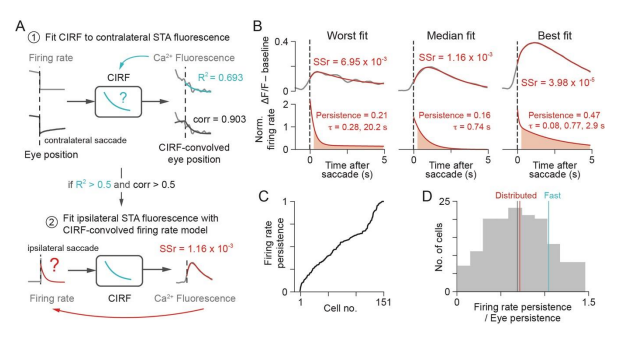
- 1081 only the fastest two time constants from the four-component plant model (cyan). (C) Same
- 1082 as B, but for larvae anaesthetized with ketamine, for which step displacements lasted 15 s or
- 1083 90 s. (D) Coefficient amplitudes and time constants for each component of the best four-
- 1084 component plant models for larvae anaesthetized with MS-222 (red, n = 10) or ketamine
- 1085 (black, n = 6). (E) Fraction of the total area under the curve (AUC) contributed by each
- 1086 component for the best four-component plant models for larvae anaesthetized with MS-222
- 1087 (red) or ketamine (black). Boxes span the 25th to 75th percentile range; whiskers show
- 1088 maximum and minimum values (MS-222, n = 10; ketamine, n = 6).





1090

1091 Figure 7 Estimating the neural drive to the oculomotor plant during fixation. (A) Eye position was deconvolved with the filter from a given larva's plant model to estimate the 1092 1093 neural drive needed to generate that eye position. (B) Perfectly stable eye position (grey) and 1094 neural drive estimates calculated from the best three-component (red) and fast two-1095 component (cvan) plant model for an example awake larva. Time courses were normalized 1096 to be 1 at 250 ms after saccade termination. (C) Same as B, but for a measured fixation. (D) 1097 Time constants of the distributed plant (black) and of the slide components of the neural drive 1098 (red circles). (E) Persistence of estimated neural drive assuming the best three- or four-1099 component (red) or the fast two-component (cyan) plant model plotted against eye position 1100 persistence. Each point represents a fixation recorded from an awake larva. Dotted line 1101 indicates a one-to-one ratio of drive persistence to eye persistence.

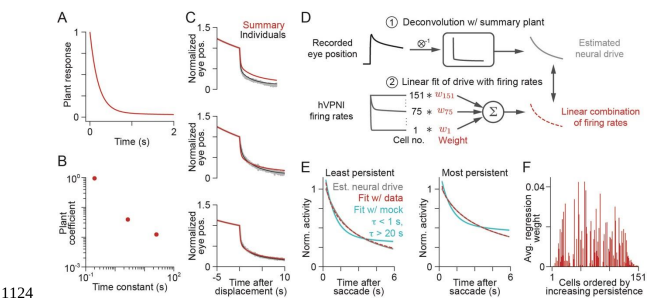


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Figure 8 The relationship between the persistence of eye position and of neuronal firing in 1104 hVPNI neurons is consistent with the neural drive required for a plant with long timescale 1105 1106 responses. (A) Illustration of the method used to recover firing rates of hVPNI neurons. First, 1107 the calcium impulse response function (CIRF) was fit to baseline subtracted saccadetriggered average (STA) Ca²⁺-sensitive fluorescence during a contralateral saccade (top). 1108 Only cells to which the CIRF fit was relatively good ($R^2 > 0.5$), and for which fluorescence 1109 was correlated to eye position when convolved with the CIRF (corr > 0.5) were included in 1110 further analysis (167/195 cells). For these cells, STA fluorescence during an ipsilateral 1111 1112 saccade was fit with a multiexponential model of post-saccadic firing convolved with the 1113 CIRF (bottom; Methods, hVPNI firing rate estimation). If the ratio of the sum of squared 1114 errors of fits to the sum of squares of ipsilateral STA fluorescence (sum of squares ratio, SSr) was greater than 0.007, the cell was excluded (16/167 cells excluded). (B) Examples of the 1115 1116 worst, median and best quality fits to ipsilateral STA fluorescence of included cells (top) and 1117 corresponding inferred firing rate functions, normalized to equal 1 at 0.25 s after saccade 1118 time (bottom). (C) Distribution of firing rate persistence (sum of red shaded areas in B) across 1119 all included neurons. (D) Histogram of the ratio of firing rate persistence to eye position 1120 persistence. Grey line indicates the median ratio. Coloured lines indicate the median ratio of

- 1121 neural drive persistence to eye persistence across larvae assuming a distributed (red) or fast
- 1122 (cyan) plant model for each larva.





1125 Figure 9 The distribution of hVPNI firing patterns is sufficient to stabilize gaze when the 1126 plant has long timescale responses. (A,B) Time course (A) and component amplitudes and 1127 time constants (B) of a single summary plant model fit to the active state step responses from 1128 all awake larvae. (C) Predicted step response given the summary plant model (red) compared 1129 to the predicted response given the best plant model for each individual larva (black). 1130 Examples are the worst (top), median (middle) and best (bottom) of the 9 reconstructed step 1131 responses, sorted by mean squared error. (D) Schematic of steps used to estimate the neural 1132 drive during fixation as a linear combination of hVPNI firing rates. Saccade-triggered 1133 average eye positions during fixation from the 6 larvae from which Ca²⁺-sensitive 1134 fluorescence was recorded were deconvolved with the summary plant model to estimate the 1135 required neural drive. For each larva we calculated a regularized linear regression of 1136 estimated neural drive onto the firing rates of all recorded hVPNI cells (n = 151 cells). (E) 1137 Grey: estimated neural drive calculated for the fish with the least (left) and most (right) 1138 persistent average eye position during fixation using the summary plant model. Red: best 1139 linear fit of hVPNI firing rates to the estimated neural drive. Cyan: band containing the best 1140 95% of linear fits to the estimated neural drive from 100 synthetic populations of mock cells 1141 whose firing rates were exponential decays with random time constants <1 s or >20 s. (F)

- 1142 Regression weight for each hVPNI cell, averaged across fits to all 6 average eye positions,
- 1143 with cells sorted by increasing firing rate persistence.

1144 DATA AVAILABILITY

- 1145 Data and code used in this study for data analysis can be found at <u>https://github.com/jay-</u>
- 1146 <u>bhasin/distributed-response-timescales</u>.
- 1147

1148 **COMPETING INTERESTS**

- 1149 The authors have no conflicts of interest to declare.
- 1150

1151 AUTHOR CONTRIBUTIONS

1152 All experiments were performed in the Tank lab at Princeton University. A.M., E.R.F.A. and 1153 D.W.T. conceived of and designed the experiments. A.M. performed the experiments and 1154 acquired the data. A.M., B.J.B., and M.S.G. conceived of and performed data analysis and 1155 modelling. A.M., B.J.B., E.R.F.A. and M.S.G. drafted the paper. All authors critically revised 1156 the paper. All authors approved the final version of the manuscript and agree to be 1157 accountable for all aspects of the work in ensuring that questions related to the accuracy or 1158 integrity of any part of the work are appropriately investigated and resolved. All persons 1159 designated as authors qualify for authorship, and all those who qualify for authorship are 1160 listed.

1161

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1169

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1320 MATHEMATICAL APPENDIX

1321 Analytical calculation of neural drive for a multiexponential plant

Here, we derive analytical expressions for the neural drive expected in the case of a linear multi-exponential plant under the inverse model formulation, extending the work of Sklavos et al. (2005). First, we provide a general formula, then specific formulae for the one- and two-exponential cases, and finally discuss simulation results for three- and four-exponential plants.

1327 Suppose that the measured eye position during normal fixations can be described by 1328 sums of n_{pos} exponential decays with time constants $\tau_{\text{pos},i} = 1/k_{\text{pos},i}$ and amplitudes $c_{\text{pos},i}$,

1329
$$y(t) = \sum_{i=1}^{n_{\text{pos}}} c_{\text{pos},i} e^{-t/\tau_{\text{pos},i}} = \sum_{i=1}^{n_{\text{pos}}} c_{\text{pos},i} e^{-k_{\text{pos},i}t}, \qquad (A.1)$$

with perfectly stable eye position represented by a single exponential decay with $k_{pos} = 0$. We assume that the eye position is generated by linear filtering of the neural drive by a plant with a multiexponential impulse response function, as in equations (1) and (2). Taking the Laplace transform of equations (1), (2) and (A.1), equation (2) becomes

1334
$$\sum_{i=1}^{n_{\text{pos},i}} \frac{c_{\text{pos},i}}{s+k_{\text{pos},i}} = Y(s) = F(s)P(s) = \left(\sum_{i=1}^{n} \frac{c_i}{s+k_i}\right)F(s),$$

1335 where $k_i = 1/\tau_i$ are the inverse time constants, as before, and Laplace transformed variables 1336 are represented with capital letters. Examining only P(s), the Laplace transform of p(t), we 1337 can turn the sum of fractions into a single fraction,

1338
$$P(s) = \frac{\sum_{i=1}^{n} c_i \prod_{j \neq i} (s+k_j)}{\prod_{i=1}^{n} (s+k_i)} = \frac{\prod_{i=1}^{n-1} (s+\lambda_i)}{\prod_{i=1}^{n} (s+k_i)},$$
(A.2)

1339 where the numerator is an n - 1 degree polynomial with roots $-\lambda_i$, and where we have used 1340 that the sum of the amplitudes c_i is 1. We can similarly write the Laplace transform of the

1341 eye position, Y(s), as a single fraction that is the ratio of two polynomials. Then, the Laplace

1342 transform of the neural drive is

1343
$$F(s) = \frac{Y(s)}{P(s)} = \frac{\prod_{i=1}^{n_{\text{pos}}-1} (s + \lambda_{\text{pos},i}) \prod_{i=1}^{n} (s + k_i)}{\prod_{i=1}^{n_{\text{pos}}} (s + k_{\text{pos},i}) \prod_{i=1}^{n-1} (s + \lambda_i)}.$$
 (A.3)

1344 Assume that $\lambda_{\text{pos},i}$, k_i , $k_{\text{pos},i}$ and λ_i are all unique. Then, by partial fraction decomposition, 1345 equation (A.3) becomes

1346
$$F(s) = 1 + \sum_{i=1}^{n_{\text{pos}}} \frac{a_{\text{pos},i}}{s + k_{\text{pos},i}} + \sum_{i=1}^{n-1} \frac{a_{\text{slide},i}}{s + \lambda_i},$$
 (A.4)

1347 and taking the inverse transform, the neural drive in the time domain is

1348
$$f(t) = \delta(t) + \sum_{i=1}^{n_{\text{pos},i}} a_{\text{pos},i} e^{-t/\tau_{\text{pos},i}} + \sum_{i=1}^{n-1} a_{\text{slide},i} e^{-\lambda_i t}, \qquad (A.5)$$

1349 which consists of a pulse, a set of components that decay as eye position, and n-11350 exponentially decaying slides with time constants $1/\lambda_i$ that may be complicated 1351 combinations of the plant coefficients and time constants, but which do not depend on the 1352 desired eye position. Note that if the desired eye position had had a component with time 1353 constant given by the plant (such that $k_{\text{pos},i} = k_j$ for some *i* and *j*), then a component with 1354 this time constant would not be required in the drive.

1355 From the residue theorem, the amplitude of any eye position component can be 1356 calculated by evaluating $(s + k_{\text{pos},i})F(s)$ at $s = -k_{\text{pos},i}$,

1357
$$a_{\text{pos},i} = [(s+k_{\text{pos},i})F(s)]_{s=-k_{\text{pos},i}} = \frac{1}{P(-k_{\text{pos},i})}.$$
 (A.6)

For the special case of perfectly stable eye position, a single component with inverse time constant $k_{pos} = 0$, this component is a step with amplitude

1360
$$a_{\text{pos}} = \frac{1}{\sum_{i=1}^{n} \frac{C_i}{k_i}} = \frac{1}{\sum_{i=1}^{n} c_i \tau_i}.$$
 (A.7)

That is, since the area contributed by each plant component *i* is $c_i \tau_i$, the amplitude of the step component is equal to the inverse of the total area of the plant filter. To provide additional intuition about the effect of long time constants on the neural drive, we below consider the simple examples of plants with either a single exponential decay or a sum of two exponential decays and for simplicity, eye position that can be described by a single exponential decay.

1366

1367 A single exponential plant

1368 Consider the case of a single exponential plant model,

$$p(t) = e^{-kt}$$

1370 Then, the Laplace transform of the neural drive is

1371
$$F(s) = \frac{s+k}{s+k_{\text{pos}}} = 1 + \frac{k-k_{\text{pos}}}{s+k_{\text{pos}}},$$

1372 and in the time domain,

1373
$$f(t) = \delta(t) + (k - k_{pos})e^{-t/\tau_{pos}},$$

where $k > k_{pos}$. Thus, in this simplest case, the neural drive consists of a pulse and a 1374 decaying component that follows eye position. In the case of stable fixation ($k_{\rm pos} = 0, \tau_{\rm pos} \rightarrow$ 1375 1376 ∞), this component is a step whose amplitude increases in proportion to the decay rate k of 1377 the plant. To interpret this result, note that convolution with an exponential filter can be 1378 thought of conceptually as a leaky integral. To generate stable eye position, an eye velocity 1379 command must be supplemented with an eye position command to compensate for the effect 1380 of the plant's leaky filter. As the time constant increases, i.e., as k decreases, the plant's 1381 integration becomes less leaky, and the step-like position command becomes less necessary. 1382

1383 A double exponential plant

1384 Next, consider a plant with two exponential components,

1385
$$p(t) = c_1 e^{-k_1 t} + c_2 e^{-k_2 t},$$

1386 with $c_1 + c_2 = 1$ and $k_1 > k_2$, which implies $\tau_1 < \tau_2$. This has Laplace transform

1387
$$P(s) = \frac{c_1}{s+k_1} + \frac{c_2}{s+k_2} = \frac{c_1(s+k_2) + c_2(s+k_1)}{(s+k_1)(s+k_2)} = \frac{s+c_1k_2 + c_2k_1}{(s+k_1)(s+k_2)}$$

1388 The required neural drive is

1389
$$F(s) = \frac{(s+k_1)(s+k_2)}{(s+k_{\text{pos}})(s+c_1k_2+c_2k_1)} = 1 + \frac{a_{\text{pos}}}{s+k_{\text{pos}}} + \frac{a_{\text{slide}}}{s+c_1k_2+c_2k_1},$$

1390 which in the time domain is

1391
$$f(t) = \delta(t) + a_{\text{pos}} e^{-t/\tau_{\text{pos}}} + a_{\text{slide}} e^{-t/\tau_{\text{slide}}},$$

1392 a combination of a pulse, a position-like, and an exponentially decaying slide component, 1393 where a_{pos} and a_{slide} are defined below. The decay rate of the slide is a linear combination 1394 of the decay rates of the two plant components,

1395
$$\frac{1}{\tau_{\text{slide}}} = k_{\text{slide}} = c_1 k_2 + c_2 k_1$$

1396 or in terms of time constants,

1397
$$\tau_{\text{slide}} = \frac{1}{c_1 k_2 + c_2 k_1} = \frac{\tau_1 \tau_2}{c_1 \tau_1 + c_2 \tau_2}$$

1398 Since $c_2 = 1 - c_1$, the slide time constant is bounded by the two plant time constants. The 1399 slide time constant is longest in the limit where the amplitude of the faster plant component 1400 is large $(c_1 \rightarrow 1, c_2 \rightarrow 0)$, in which case it approaches τ_2 , the time constant of the slower 1401 plant component. It is shortest in the limit in which the amplitude of the faster plant 1402 component is very small $(c_1 \rightarrow 0, c_2 \rightarrow 1)$, in which case it approaches τ_1 .

1403 The purpose of the slide can be seen clearly in the case of perfectly stable desired eye 1404 position, $k_{pos} = 0$. Applying neural drive with only a pulse and position (step) component to 1405 the plant, the eye position due to each plant component *i* is

1406
$$c_{i}e^{-k_{i}t} * (\delta(t) + a_{pos}u(t)) = a_{pos}\frac{c_{i}}{k_{i}} + c_{i}\left(1 - \frac{a_{pos}}{k_{i}}\right)e^{-k_{i}t},$$

1407 where u(t) is the Heaviside or unit step function. We can find a step amplitude a_{pos} such 1408 that the exponential decay term in the above expression is cancelled for each plant component 1409 individually, but we will be unable to simultaneously cancel the corresponding exponential 1410 decay resulting from the other plant component (a pulse/step mismatch). This is corrected by 1411 adding the additional slide component to the drive.

1412The amplitudes of the position and slide components of the drive are respectively1413given by

1414
$$a_{\rm pos} = \frac{(k_1 - k_{\rm pos})(k_2 - k_{\rm pos})}{k_{\rm slide} - k_{\rm pos}}$$

1415
$$a_{\text{slide}} = \frac{(k_1 - k_{\text{slide}})(k_{\text{slide}} - k_2)}{k_{\text{slide}} - k_{\text{pos}}}$$
$$= \frac{c_1 c_2 (k_1 - k_2)^2}{k_{\text{slide}} - k_{\text{pos}}}.$$

1416 For simplicity, let us consider perfectly stable fixations, $k_{pos} = 0$, so that again the 1417 eye position component is a step. Then, the ratio of the amplitude of the slide to the amplitude 1418 of the step is given by

1419
$$\frac{a_{\text{slide}}}{a_{\text{pos}}} = \frac{c_1 c_2 (k_1 - k_2)^2}{k_1 k_2}$$

For a fixed choice of time constants, the ratio is proportional to the product $c_1c_2 = c_1(1 - c_1)$. If either c_1 or c_2 is sufficiently close to 1, then the amplitude of the slide will be extremely small compared to the step. The ratio is maximized for $c_1 = c_2 = 1/2$.

1423 In this study, we encountered the situation in which plant components contributed 1424 similar areas to the plant, an important parameter setting we will now consider. In the case 1425 of two equal area components, with $c_1/k_1 = c_2/k_2$, the coefficients of the plant components 1426 become

1427
$$c_{1} = \frac{k_{1}}{k_{1} + k_{2}}$$
$$c_{2} = \frac{k_{2}}{k_{1} + k_{2}}.$$

1428 Then, the time constant of the slide is the average of the two plant time constants,

1429
$$au_{slide} = \frac{1}{2}(\tau_1 + \tau_2),$$

1430 and the amplitudes of the step and the slide are

1431
$$a_{\text{pos}} = \frac{1}{2}(k_1 + k_2)$$
$$a_{\text{slide}} = \frac{1}{2}\frac{(k_1 - k_2)^2}{k_1 + k_2}$$

In this case, the contribution of the slide is appreciable only if the inverse time constants of the two plant components are far apart. For the equal areas case, we also can understand easily the effect of a long time constant on the required neural drive. For a fixed choice of time constant τ_1 and amplitude c_1 for the faster component, as the time constant of the slower component becomes longer, the slide time constant increases, the amplitude of the step decreases, and the amplitude of the slide increases.

1438 Three- and four-component plants

Using numerical simulations, we extended this analysis to three- and four-component plants. In each case, we generated 1 million plants with coefficients and inverse time constants chosen uniformly at random between 0 and 1, with coefficients then normalized so that they sum to 1. We calculated the time constants of the slide components by solving for the zeros $-\lambda_i$ of the Laplace transform of each plant (equation (A.2)). We found that, in all cases, there was exactly one slide time constant between each pair of neighbouring plant time constants.