# 1 Central vestibular tuning arises from patterned convergence of otolith afferents

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# 12 Abstract:

- 13 As sensory information moves through the brain, higher-order areas exhibit more
- 14 complex tuning than lower areas. Though models predict this complexity is due to
- 15 convergent inputs from neurons with diverse response properties, in most vertebrate
- 16 systems convergence has only been inferred rather than tested directly. Here we
- 17 measure sensory computations in zebrafish vestibular neurons across multiple axes in
- *vivo*. We establish that whole-cell physiological recordings reveal tuning of individual
- 19 vestibular afferent inputs and their postsynaptic targets. An independent approach,
- 20 serial section electron microscopy, supports the inferred connectivity. We find that
- afferents with similar or differing preferred directions converge on central vestibular
- neurons, conferring more simple or complex tuning, respectively. Our data also resolve
   a long-standing contradiction between anatomical and physiological analyses by
- revealing that sensory responses are produced by sparse but powerful inputs from
- vestibular afferents. Together these results provide a direct, quantifiable demonstration
- 26 of feedforward input convergence in vivo.
- 27 **Key words:** neural computation; sensory encoding; feedforward excitation;
- vestibulospinal neuron; electrical synapse; high-pass tuning; sensorimotor
- 29 transformation; body balance

#### 30 Introduction

Neurons compute information from many different synaptic inputs. A central 31 32 challenge in understanding neuronal circuits is determining how the tuning and 33 connectivity of these inputs affect the resulting computations. For example, neurons in 34 visual cortex exhibit simple or complex orientation tuning, which is thought to derive from the convergence of presynaptic inputs with distinct tuning properties (Hubel and 35 Wiesel, 1962, Alonso and Martinez, 1998). Computational models of such input-output 36 37 relationships have fundamentally shaped the way we think of information processing in the brain (Felleman and Van Essen, 1991, LeCun et al., 2015). However, these models 38 generally require assumptions about many parameters that can only be measured with 39 40 incompatible approaches: the tuning of the presynaptic population, input connectivity, and synaptic strengths, as well as the activity of the postsynaptic neuron itself. Direct 41 measurements of these parameters simultaneously are prohibitively difficult in most 42 systems, making it hard to define neuronal computations in vivo. 43

44 Vestibulospinal (VS) brainstem neurons receive direct vestibular sensory inputs from peripheral vestibular afferents (Boyle et al., 1992) and project to the spinal cord 45 (Boyle and Johanson, 2003). Understanding the neuronal computations of VS neurons 46 47 would not only inform how vestibular sensory signals are processed in the brain, but 48 also provide a mechanistic view of sensorimotor transformation. VS neurons, like other 49 central vestibular neurons, produce diverse responses to head movement. During head tilt or acceleration, some central vestibular neurons exhibit simple cosine-tuned 50 51 responses, similar to those of the afferents: the strongest activity is evoked by movements in a preferred direction, with little or no response in the orthogonal direction. 52 In contrast, other central vestibular neurons exhibit more complex responses, including 53 bidirectional responses (Peterson, 1970) and spatiotemporally complex tuning (Angelaki 54 55 et al., 1993). A vectorial model predicts that convergence of several simple cosine-56 tuned afferents can fully account for the response of either a simple or a complex 57 central vestibular neuron, depending on whether those afferents are similarly tuned or differently tuned (Angelaki, 1992). However, as in other systems, this model has been 58 59 technically challenging to test experimentally.

We chose to address this question in the small brain of the larval zebrafish. The VS circuit was previously identified in the larval zebrafish as the homolog of mammals (Kimmel et al., 1982), which becomes functional as early as 3 days post fertilization (dpf) (Mo et al., 2010). The accessibility of the larval zebrafish brain for intracellular recording from identified VS neurons allows us to investigate how central vestibular neurons compute sensory signals in vertebrates.

66 Here we establish a novel approach to record sensory evoked responses *in vivo* 67 from VS neurons in the larval zebrafish. We find that individual afferents evoke large 68 amplitude-invariant excitatory postsynaptic currents (EPSCs), allowing us to separate 69 distinct afferent inputs that converge onto a given VS neuron. This provides a 70 mechanism to simultaneously measure the sensory tuning and synaptic strength of 71 each converging afferent, as well as the response of the postsynaptic neuron. We show

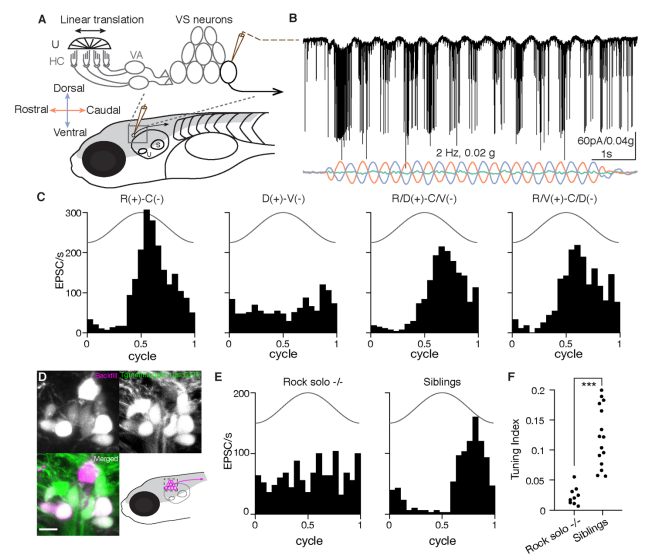


Figure 1: Sensory-evoked responses *in vivo* in vestibulospinal (VS) neurons A. Schematic representing *in vivo* patch clamp recording configuration and vestibular afferent circuit in larval zebrafish, U: utricle, HC: hair cells, VA: vestibular afferents, S: saccule.

B. Example recording trace from a VS neuron in voltage clamp mode during 2 Hz, 0.02 g translational movement on the R/V(+)-C/D(-) axis. EPSCs, black; acceleration in three body axes, colored (orange, rostral[+]-caudal[-]; green, dorsal[+]-ventral[-]; blue, ipsilateral[+]-contralateral[-]).

C. Sensory-evoked EPSC responses to translation in four different axes for the same VS neuron as in B, across 12 cycles. Solid line, acceleration (2 Hz, 0.02 g).

D. Tg(nefma:gal4; UAS:GFP) (green) colabels VS neurons identified by dye backfilling (magenta) from spinal cord. Scale bar:  $5 \,\mu$ m

E. Sensory responses of a VS neuron in the best direction in a rock solo -/- (left) and in a het/WT sibling (right).

F. Summary of tuning index in the best direction for all VS neurons recorded in rock solo -/- (9 neurons, 5 fish) and siblings (15 neurons, 10 fish). Mann-Whitney U test, p=6.5e-5

- that afferents with similar tuning direction preferentially converge, producing simple 72
- 73 tuning in the VS neuron. Furthermore, the smaller number of cells with complex
- 74 bidirectional responses receive input from differently tuned afferents, with consequent
- 75 simple or complex spiking. We also show that these afferent inputs are sufficient to
- 76 predict the tuning of the VS neuron. Together, this work reveals how central neurons in
- the brain compute sensory information from their presynaptic inputs. 77

#### 78 **Results:**

#### 79 Sensory evoked responses of vestibulospinal neurons in vivo

80 Traditionally, measurements of neuronal responses to vestibular stimuli have been accomplished by unit recordings (Angelaki and Dickman, 2000, Schor et al., 1984, 81

- 82 Fernandez and Goldberg, 1976a). Directly measuring vestibular-evoked synaptic
- currents in central neurons in vivo has been technically challenging (Arenz et al., 2008, 83
- 84 Chabrol et al., 2015). We designed a custom whole-cell electrophysiology rig to deliver
- translational motion stimuli to 4-7 dpf larval zebrafish via an air-bearing motorized sled
- 85 (Fig. 1A). This setup allows intracellular measurement of sensory-evoked responses 86
- from vestibulospinal (VS) neurons on multiple axes in vivo for the first time, to the best 87
- 88 of our knowledge. To target identified VS neurons, we generated a Tg(nefma:gal4;
- UAS:GFP) line, whose labelling overlaps dye backfilling (Figs.1D, S1) from the spinal 89
- 90 cord, consistent with evidence of *Nefm* expression in mammalian vestibular neurons
- 91 (Kodama et al., 2020). We recorded spontaneous EPSCs in voltage clamp, at overall
- rates varying from 1 to 365 EPSC/s. Delivery of translational movement evoked 92
- 93 corresponding modulations in EPSC frequency (Fig. 1B). The extent of modulation
- varied depending on the direction of the stimulus delivered across four different axes 94
- (Fig. 1C). In this example neuron, EPSC rate was modulated most strongly in the 95
- rostral-caudal (R-C) axis and weakly in the dorsal-ventral (D-V) axis, with intermediate 96
- 97 strength responses for the diagonal directions (R/D-C/V and R/V-C/D).
- 98 Response to translational movement could derive from the vestibular or other sensory inputs. In larval zebrafish, the anterior otolith (utricle) is the sole functional vestibular 99
- 100 sensor (Riley and Moorman, 2000). To examine whether utricular signaling is necessary
- 101 for the observed tuning, we measured the sensory response of VS neurons in the
- otogc.1522+2T>A -/- (rock solo) animals, which lack the utricle (Mo et al., 2010, Roberts et 102
- al., 2017). Translational stimuli were ineffective at modulating EPSC rate in VS neurons 103
- of rock solo homozygotes in contrast to wild-type/heterozygous siblings (representative 104
- examples, Fig. 1E). Across all recordings, VS neurons of rock solo -/- animals exhibit 105
- largely untuned EPSCs compared to siblings, as guantified by a tuning index (Fig. 1F). 106
- 107 Thus, this approach reveals sensory-evoked synaptic responses encoding directional
- vestibular stimuli in identified VS neurons in vivo. 108

#### 109 Mixed electrical and chemical synapses mediate the transmission from otolith afferents to VS neurons 110

What properties define the vestibular afferent synapse onto VS neurons? In 111 rodents, vestibular afferent synapses onto vestibulo-ocular reflex neurons exhibit 112

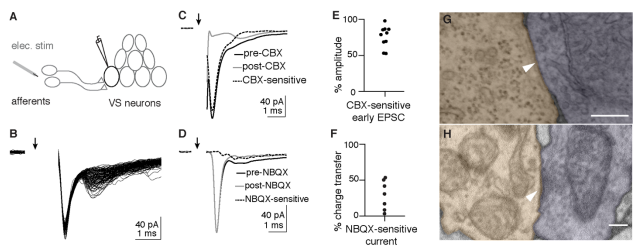


Figure 2: Otolith afferent to VS neuron transmission is mediated by mixed electrical and chemical synapses

A. Schematic of whole-cell recording configuration from VS neuron while electrically stimulating otolith afferents.

B. Example EPSCs evoked by electrical stimulation of the otolith afferents; 105 EPSCs overlaid. Arrow indicates onset of stimulation. Stimulus artifact is blanked.

C. Carbenoxolone (CBX) diminishes the fast component of evoked EPSCs

D. NBQX abolishes the second, slower component of evoked EPSCs

E. Group data quantifying the reduction of early EPSC amplitude by CBX

F. Group data quantifying the total charge transfer that is abolished by NBQX application

G. Example EM image of gap junction between identified otolith afferent (pseudocolored purple) and VS neuron (orange). Scale bar: 200 nm.

H. Example EM image of chemical synapse between otolith afferent (purple) and VS neuron (orange). Scale bar: 200 nm.

amplitude-invariant synaptic transmission, mediated by specialized vesicular release 113 114 machinery (Bagnall et al., 2008, McElvain et al., 2015, Turecek et al., 2017). To 115 characterize afferent synaptic input to VS neurons, we electrically stimulated the 116 vestibular (anterior statoacoustic) ganglion while recording from VS neurons in voltage 117 clamp (Fig. 2A). Stimulation evoked a synaptic current with two components. The first component had fast kinetics with short latency ( $0.56 \pm 0.28$  ms, n=8), low jitter (0.05118  $\pm$  0.04 ms, n=8), and invariant EPSC amplitude (SD: 6.7 $\pm$ 3.9%, normalized to peak) 119 120 across trials. In contrast, the second component had slower kinetics and variable 121 amplitudes (Fig. 2B). We dissected the two components of evoked EPSCs 122 pharmacologically. Bath application of the gap junction blocker carbenoxolone (CBX. 123 500  $\mu$ M) during afferent stimulation substantially reduced the first component of the EPSC (Figs. 2C, E). In contrast, bath application of the AMPA receptor antagonist 124 125 NBQX (10  $\mu$ M) abolished the second component of synaptic current (Fig. 2D). Furthermore, the fast EPSCs were not reversed by changing the holding potential (Fig. 126 127 S2), a signature behavior of electrical synaptic transmission (Akrouh and 128 Kerschensteiner, 2013). Thus, the early and late components of afferent-evoked synaptic currents are mediated by gap junctions and AMPA receptors, respectively. 129 Across VS neurons, the NBQX-sensitive currents accounted for 27.1±20.2% of total 130 charge transfer (n=7, Fig. 2F), demonstrating that gap junctional current is the major 131

132 component mediating synaptic transmission.

133 To evaluate ultrastructural evidence for mixed synaptic transmission, we re-134 imaged existing serial ultrathin sections of a 5.5 dpf larval zebrafish (Hildebrand et al., 135 2017) at sufficiently high resolution (1-4 nm/px) to identify synaptic contacts between myelinated utricular afferents and VS neurons, identified anatomically. We found both 136 137 tight junction structures (Fig. 2G), and vesicles apposed to a postsynaptic density (Fig. 2H) at appositions between utricular afferent and VS neurons, consistent with 138 139 anatomical evidence for mixed electrical / chemical transmission at this synapse in adult fish (Korn et al., 1977) and rat (Nagy et al., 2013). Together, these results demonstrate 140 141 that VS neurons receive vestibular afferent inputs mediated by amplitude-invariant gap junctional (electrical) and variable amplitude glutamatergic (chemical) synapses. 142

# 143 Inferring afferent tuning from distinct EPSCs

144 Because electrically mediated EPSCs from afferents to VS neurons exhibited a 145 fixed amplitude, we hypothesized that we could distinguish the activity of individual otolith afferents converging onto a given VS neuron by their characteristic EPSC 146 147 amplitudes. Indeed, spontaneous and sensory-evoked EPSCs recorded in VS neurons often fell into distinctive size bins, as visualized in a histogram of EPSC amplitudes 148 (Figs. 3A, B). EPSCs were sorted into three clusters with unsupervised learning (see 149 Methods), primarily leveraging their amplitudes. Each of these EPSC clusters showed a 150 151 stereotypical amplitude and waveform in this example neuron (Fig. 3A, inset). To test 152 whether each cluster of EPSCs amplitudes corresponds to an individual afferent, we 153 used an approach derived from spike sorting: temporal autocorrelation to test for 154 refractory periods within EPSC event times. Physiologically, one afferent cannot generate two action potentials within its refractory period (Fernandez et al., 1972); thus 155

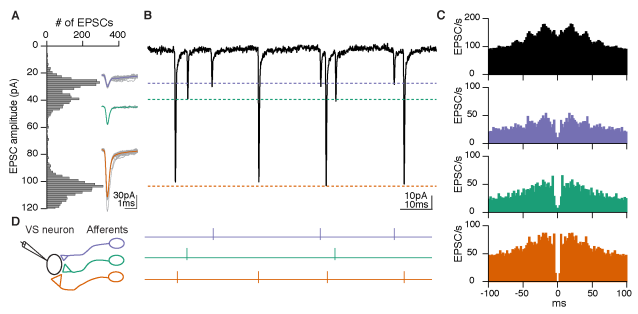


Figure 3: Distinct EPSC amplitudes reflect individual afferent inputs

A. Histogram of spontaneous and sensory-evoked EPSC amplitude distribution of the same VS neuron as Fig. 1B. Inset, overlay of individual EPSCs (gray) and average (colored) for each amplitude bin.

B. Example trace of EPSCs exhibiting stereotypic shapes and amplitudes in three clusters, corresponding to each amplitude bin in A.

C. Auto-correlogram of all EPSCs recorded from the VS neuron (top, black) or divided into three clusters based on EPSC amplitudes (bottom, colored). A refractory period around 0 ms only occurs for EPSCs within each cluster, but not across all EPSCs.
D. Schematic of three different otolith afferents converging onto one VS neuron, each eliciting EPSCs with a distinct amplitude (represented by different synaptic sizes). Right, spike activities of three afferents inferred from B.

the EPSCs elicited by that afferent should exhibit a refractory period as well. An auto-156 157 correlogram of all EPSC event times in this example neuron did not display a refractory 158 period (Fig. 3C, top). In contrast, an auto-correlogram within each EPSC cluster 159 exhibited a clear refractory period around 0 ms (Fig. 3C, bottom). Furthermore, cross-160 correlograms between EPSC clusters did not show this structure, consistent with the notion that they arise from independent inputs (Fig. S3). Accordingly, we can interpret 161 these three EPSC clusters as deriving from the activity of three distinct presynaptic 162 163 afferents (Fig. 3D, left). Because of the high fidelity of electrical transmission, each 164 EPSC cluster effectively reads out the spiking of an individual afferent, allowing us to 165 measure presynaptic activity via postsynaptic recording (Fig. 3D, right).

166 To test this interpretation of electrophysiological data with a completely 167 independent approach, we reconstructed the whole volume of myelinated utricular inputs onto 11 VS neurons from a high resolution re-imaged serial section EM dataset 168 acquired from the right side of one 5.5 dpf larval zebrafish (Fig. 4A, B). We found that 169 170 the connection between myelinated utricular afferents and VS neurons was relatively 171 sparse. All VS neurons were contacted by at least two utricular afferents, but some afferents did not innervate any VS neurons (Fig. 4C). These reconstructions showed 172 173 that a range of 2-6 afferents (mean+std: 3.4+1.4) converged onto each VS neuron (Fig. 174 4D). We compared these numbers to those derived from whole-cell physiology, where we inferred the number of convergent afferents from the number of EPSC clusters. 175 176 Across all VS neuron recordings, we found a range of 0-5 afferents (1.7±1.3) converged onto each VS neuron (Fig. 4E). The result from anatomical reconstruction is largely 177 consistent with the overall distribution of afferent contacts as measured by whole-cell 178 179 physiology, presumably with some small-amplitude EPSCs elicited by the afferents not 180 successfully clustered. Therefore, these results demonstrate that synaptic inputs from individual vestibular afferents can be separated by their stereotypic EPSC waveforms, 181 yielding inferred afferent convergence consistent with high-resolution anatomical 182 183 connectivity.

# 184 Spatial tuning of inferred otolith afferents

By recording from one VS neuron, we can infer the activity of its presynaptic 185 afferents. This approach thus offers a unique opportunity to measure the sensory tuning 186 187 of several convergent afferents simultaneously. To determine the spatial tuning of 188 convergent afferent inputs, we delivered 2 Hz, +0.02 g sinusoidal translational stimuli 189 on four axes in the horizontal plane and recorded the sensory-evoked EPSCs, as 190 shown for an example VS neuron (Fig. 5A). In this example neuron, the inferred utricular afferent (EPSC cluster) with the largest synaptic amplitude responded best to 191 caudally-directed acceleration, while two others responded with varying sensitivities to 192 rostrally-directed acceleration, in all cases with phase leads relative to peak 193 194 acceleration (Fig. 5B). With these measurements, we can derive the preferred tuning 195 direction, gain and phase of each afferent, as represented by the direction and length of 196 a vector (Fig. 5B, right). To validate the consistency of the vectorial representation, we 197 used a previously established approach (Schor et al., 1984) to quantify the tuning vectors with separately measured responses to two circular stimuli (Fig. S4 B), which 198

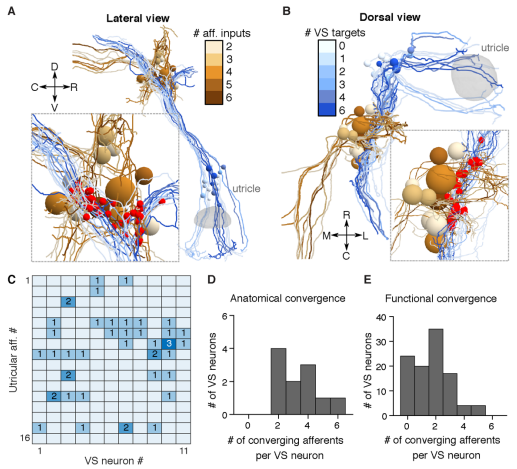


Figure 4: Anatomical reconstruction reveal similar convergence pattern as physiology A. Serial-section EM reconstruction (lateral view) of all myelinated utricular afferents (blues) and VS neurons (browns) on the right side of one animal (5.5 dpf). Inset, identified synaptic contacts between afferents and VS neurons (red). Color scale represents number of distinct afferents synapsing with a given VS neuron (browns). VS neurons with greater number of afferent inputs are located more ventrally.

B. Dorsal view of the same reconstruction as A. Color scale represents number of VS neurons contacted by a given afferent (blues).

C. Number of distinct synaptic contacts from each utricular afferent onto each VS neuron, based on serial-section EM reconstruction.

D. Histogram of the numbers of distinct afferents converging onto each VS neuron, as measured by serial-section EM reconstruction (11 neurons, 1 fish)

E. Histogram of the numbers of distinct afferents converging onto each VS neuron, as inferred from whole-cell physiology recording (104 neurons, 89 fish)

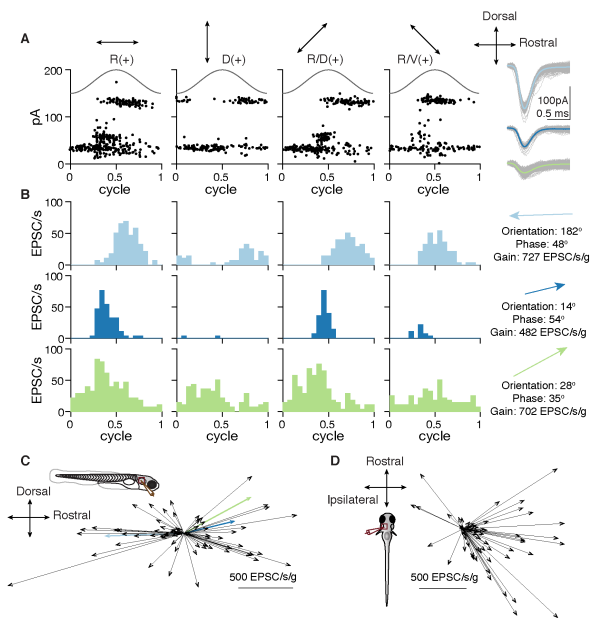


Figure 5: Spatial tuning of inferred otolith afferents

A. EPSC responses of an example neuron in response to 2 Hz, 0.02 g translational stimuli (solid sinusoidal line, acceleration) on 4 different axes (top, arrows). Each dot represents one EPSC; note three EPSC clusters with distinct amplitudes. Right, overlay of individual EPSCs (gray) and average (colored) for each cluster.

B. EPSC tuning of three clusters. Right, vectors representing the maximum tuning direction, phase, and gain of each inferred afferent corresponding to an EPSC cluster.
C. Maximum tuning directions of all afferents from VS neurons recorded from fish oriented side-up. Colored arrows represent tuning of afferents in B (69 afferents, 43 neurons, 33 fish)

D. Maximum tuning directions of all inferred afferents from VS neurons recorded from fish oriented dorsal-up. (60 afferents, 36 neurons, 36 fish)

showed similar preferred directions as those measured by translational stimuli (Figs. S4 199 200 A-C). Across all recordings with the animal oriented side-up, tuning of afferents was 201 strong in the rostral (30/69) and caudal (31/69) directions, but relatively weak in the 202 dorsal (4/69) and ventral (4/69) directions, as represented by an overlay of all inferred 203 afferent tuning vectors (Fig. 5C). When fish were oriented dorsal-up, the axes tested 204 were rostral-caudal and ipsilateral-contralateral (motion along an axis from one ear to the other). In this position, most afferents were strongly tuned to acceleration towards 205 206 the contralateral direction (31/60), some exhibited preferential tuning to the acceleration to the rostral (4/60) and caudal (20/60) directions, and only 5/60 afferents were tuned to 207 208 the ipsilateral direction (Fig. 5D). These results showed that each afferent in the larval zebrafish exhibits selective responses to different translational stimuli. Afferents overall 209 responded best to acceleration towards the contralateral, rostral and caudal directions, 210 which correspond to ipsilateral, nose-up and nose-down tilts in postural change 211 (Angelaki and Cullen, 2008), consistent with the distribution of hair cell polarity in the 212

213 utricular macula (Haddon et al., 1999).

# 214 Temporal tuning of inferred otolith afferents

215 The sensitivity and phase of vestibular afferents varies for motion at different frequencies (Fernandez and Goldberg, 1976b). The tuning of otolith afferents ranges 216 from typically more jerk-encoding (derivative of acceleration) at low frequencies to more 217 acceleration-encoding at high frequencies. What temporal tuning profile do afferents in 218 219 larval zebrafish exhibit? We applied translational stimuli with different frequencies (0.5-8 220 Hz,  $\pm 0.02$  g) on the rostral-caudal axis. In the example neuron, all three inferred otolith 221 afferents showed similar tuning, with progressively stronger responses with increasing 222 frequencies of stimulation (Fig. 6A). Across group data acquired at both  $\pm 0.02$  g and 223  $\pm 0.06$  g, the average tuning gain increased 3.3-fold (0.02 g) and 2.3-fold (0.06 g) from 0.5 Hz to 8 Hz (Fig. 6B). Most afferents (39/48) showed at least 2-fold increase from 0.5 224 225 Hz to 4 Hz in tuning gain at either 0.02 g or 0.06 g. Only one afferent had relatively flat 226 gain (< 50% increase) at both 0.02 g and 0.06 g, and its tuning was overall weak (mean gain: 1.88 and 2.24 EPSC/s respectively), suggesting it was less sensitive or not tuned 227 on the rostral-caudal axis. Regardless of tuning direction (rostral: 44%, 21/48; caudal: 228 56%, 27/48), afferents exhibited a phase lead relative to peak acceleration at various 229 230 tested stimulus magnitudes and frequencies ((Fig. 6C, S5). On average, the phase lead at low frequency (0.5 Hz) was 84.0° for 0.02 g and 78.6° for 0.06 g. At high frequency (8 231 232 Hz), the phase lead was reduced to 33.6° for 0.02 g and 39.3° for 0.06 g. The temporal 233 dynamics of the afferents resembled those of previously reported irregular units 234 (Goldberg et al., 1990), with low spontaneous firing rates (10.28+9.1 EPSC/s) and larger coefficients of variation (CV). The average CV across inferred afferents was 235 236 0.97±0.24, and the smallest CV was 0.5 (Fig. S6), indicating that no regular-firing otolith afferents were detected synapsing onto VS neurons. We conclude that the otolith 237 238 afferents act as a high-pass filter, encoding a mixture of acceleration and jerk, similar to 239 otolith afferents in primates (Laurens et al., 2017).

# 240 **Preferential convergence**

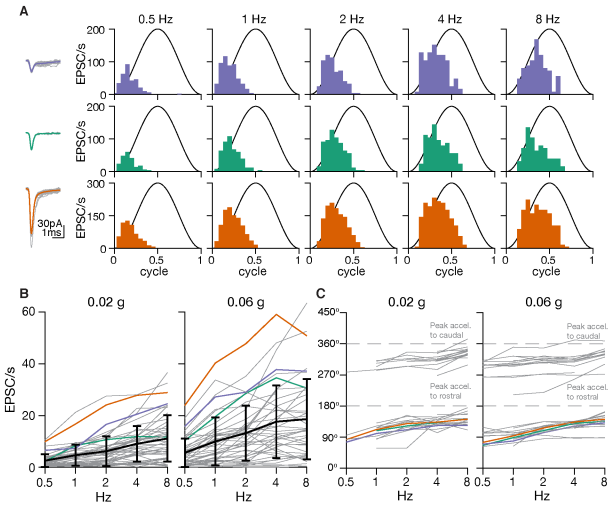


Figure 6: Temporal tuning of inferred otolith afferents

A. Sensory tuning of afferent inputs to one VS neuron during translational movement at 5 different frequencies in the rostral(+)-caudal(-) axis. Left, EPSC waveforms of three different clusters recorded from one VS neuron. Right, temporal tuning profile of each EPSC cluster on the rostral-caudal axis.

B. Gains of inferred afferents across different frequencies of translational acceleration. Gray, individual afferents; colored, afferents from A; black, mean and standard deviation of gains from all afferents (0.02 g, 48 afferents; 0.06 g: 46 afferents; 25 neurons, 20 fish)

C. Phases of inferred afferents across frequencies, relative to sinusoidal stimulus.  $180^{\circ}$  (0.5 cycle in A) represents the peak of acceleration towards rostral direction;  $360^{\circ}$  represents the peak of acceleration towards caudal direction (0 or 1 cycle in A). Data were thresholded to only include afferents whose gain was > 5 EPSC/s (0.02 g, 36 afferents; 0.06 g, 38 afferents; 25 neurons, 20 fish)

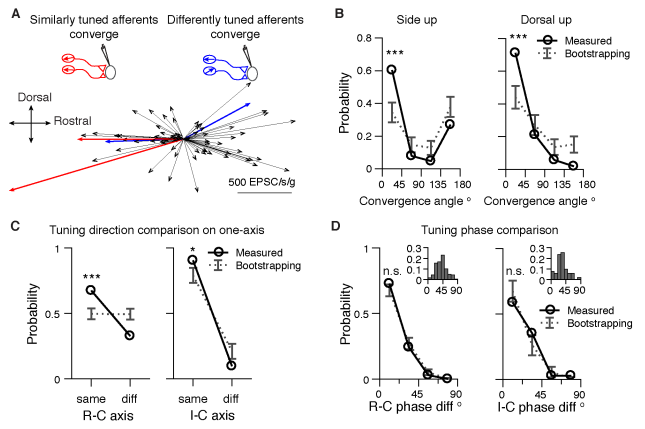


Figure 7: Afferents with similar tuning direction preferentially converge

A. Example of two pairs of converging afferents from two VS neurons in side-up fish. Red, converging afferents are similarly tuned, with small convergent angle between the pair; Blue: converging afferents are differently tuned, with large convergent angle between the pair.

B. Probability distribution of converging angles for measured and randomly generated afferents pairs in side up fish and dorsal up fish. Two tailed z-test, side up, 0-45°: p=2e-5, 135-180°: p=0.08. (63 afferent pairs); dorsal up, 0-45°: p=8e-5, 135-180°: p=0.007. (52 afferent pairs)

C. Probability distribution of converging afferents tuned to the same direction vs different direction, on the rostral-caudal and ipsilateral-contralateral axis. Two tailed z-test, R-C, same: p=1e-5, diff: p=4e-5 (150 afferent pairs); I-C, same: p=0.044, diff: p=0.044 (52 afferent pairs).

D. Probability distribution of phase difference for converging afferents, on the rostralcaudal and ipsilateral-contralateral axis. Two tailed z-test, R-C, 0-22.5° p=0.26 (103 afferent pairs); I-C, 0-22.5°, p=0.28 (34 afferent pairs). Inset: distribution of tuning phase of afferents, R-C, 177 afferents; I-C, 60 afferents; 90° represents the peak of acceleration of preferred direction (2 Hz, 0.02 g).

Individual VS neurons can receive inputs from afferents with similar (Fig. 6A) or 241 242 different tuning (Fig. 5B). Is afferent tuning convergence random or structured? The 243 responses of inferred afferents that converge onto the same VS neuron were 244 represented by their tuning vectors (Fig. 7A). The angle between the vectors indicates 245 the similarity of convergent inputs, with a small angle for a VS neuron with similarly 246 tuned inputs and a large angle for a VS neuron with differently tuned inputs. From 43 247 VS neurons recorded in the side-up orientation, 60% (38/63) of converging afferent 248 pairs had small angles (<45°) and 27% (17/63) had large angles (>135°). Compared to 249 a random pairing angle distribution generated by bootstrapping, the percentage of 250 similarly tuned convergent afferent pairs was significantly higher than chance (Fig. 7B, left). From 36 VS neurons recorded in the dorsal-up orientation, there were 71% (37/52) 251 of inferred afferent pairs with a converging angle smaller than 45°, and only 2% (1/52) 252 with a converging angle larger than 135° due to the small number of ipsilaterally tuned 253 afferents (Fig. 5D). Nonetheless, the probability of similarly tuned afferent convergence 254 255 (<45°) was significantly higher than that chance (Fig. 7B, right). For afferent pairs with converging angle larger than 45° (45°-90°, 90°-135°, 135°-180°), their probabilities was 256 257 slightly lower than their respective estimated distribution by bootstrapping. Accordingly, on a given body axis (R-C or I-C), convergent afferents are also more likely to encode 258 259 similar tuning directions (Fig. 7C). These results suggest that afferents with similar 260 tuning direction preferentially converge at rates exceeding what would be expected by 261 random connectivity.

262 Do converging afferents also have similar tuning phase regardless of their tuning direction? Most afferents are phase-leading with 2 Hz, 0.02 g stimulation (Figs. 6 and 263 7D, inset), and the phase difference between afferents is small (R-C:  $41^{\circ} \pm 16^{\circ}$ , n=177, 264 265 I-C: 33° ±17°, n=60). Consequently, most afferent pairs (R-C, 68±4.6%; I-C, 68±8%) selected randomly have very small phase difference (phase diff. < 22.5°) (Fig. 7D). Both 266 267 the probability of converging afferents having similar phase (phase diff. < 22.5°) (R-C, 73%, 75/103; I-C, 59%, 20/34) and the cumulative distribution (Fig. S7) lay within the 268 269 bootstrap predications on the rostral-caudal and ipsilateral-contralateral axes. 270 Therefore, tuning phase between converging afferent pairs is similar, in accordance with 271 their relatively homogeneous distribution.

In conclusion, we found that afferents forming synaptic connections with the
same postsynaptic VS partner typically have similar spatiotemporal tuning properties. In
particular, afferents with similar tuning direction preferentially converge, which explains
the long-standing observation that most VS neurons exhibit simple cosine tuning
(Peterson, 1970, Schor et al., 1984). However, a non-negligible number of VS neurons
receive convergent input from differently tuned afferents, a potential source for the
complex spatiotemporal tuning of central vestibular neurons.

# 279 **Complexity of central tuning is determined by the similarity of afferent inputs.**

Complex sensory tuning of central neurons is thought to arise from convergence of more simply tuned inputs with differing spatial and temporal properties, in vestibular (Angelaki et al., 1993), as well as visual (Jia et al., 2010) and somatosensory (Petersen, 2007, Roy et al., 2011) systems. For example, complex tuning such as bidirectional

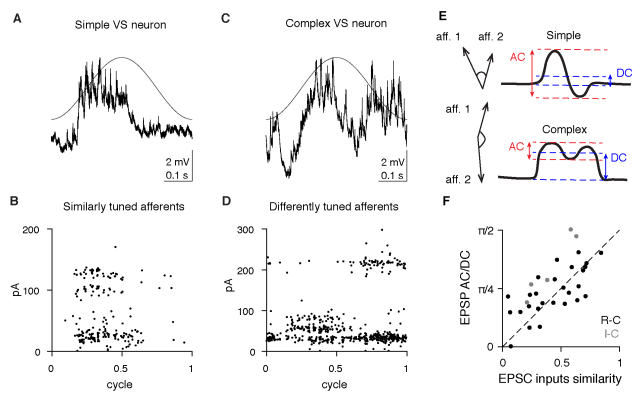


Figure 8: Subthreshold tuning responses of VS neurons are explained by the similarity of tuning of afferent inputs

A. Average membrane potential change in a VS neuron with simple response to 2 Hz, 0.02 g translational movement on the rostral(+)-caudal(-) axis.

B. EPSC responses for the simple cell shown in A. All inferred afferents exhibit similar tuning.

C. Average membrane potential change in a VS neuron with complex subthreshold response to 2 Hz, 0.02 g translational movement on the rostral(+)-caudal(-) axis

D. EPSC responses for the complex cell shown in B. Different inferred afferents exhibit distinct tuning, as shown by the different temporal responses of the large amplitude and medium amplitude EPSCs.

E. Schematic of different magnitudes of AC and DC responses in a simple and complex cell. Left, the hypothesized models of similarly tuned or differently tuned convergent afferent inputs underlying simple or complex responses, respectively.

F. Correlation of EPSC inputs similarity index and EPSP AC/DC response ratio (see Methods), for all non-spiking VS neurons with multiple convergent afferents. Sensory tuning of afferent inputs and EPSPs was measured on the R-C axis (black, n=27) and I-C axis (grey, n=5). Dashed, unity line. Pearson's R: 0.67

(Peterson, 1970) and broadly tuned sensory responses (Angelaki, 1992) of central
vestibular neurons can be computationally reconstructed from multiple modelled cosinetuned inputs. However, directly measuring these inputs has been technically difficult,
and it is unclear whether such models can sufficiently explain the activity of central
neurons. Therefore, we took advantage of the inferred afferent spiking to examine
whether the tuning of VS neurons can be constructed from the convergence of otolith
afferents.

291 We observed that different VS neurons showed simple or complex membrane potential responses to translational stimuli on the rostral-caudal axis. An example 292 293 simple cell was only depolarized during a specific phase of acceleration (Fig. 8A). 294 whereas a complex cell exhibited multiple depolarized periods during the stimulus (Fig. 295 8C). Next, we measured the EPSC tuning in the same VS neurons. In the example 296 simple neuron, sensory evoked EPSCs exhibit three distinct amplitudes (Fig. 8B), 297 indicating three afferents converge onto the cell. These three afferents showed similar 298 tuning to each other, with strongest responses for rostrally-directed acceleration. In 299 contrast, the four inferred afferents that converge onto the example complex cell exhibited a different tuning pattern. Two afferents were tuned to rostrally-directed 300 301 acceleration and the other two to caudally-directed acceleration (Fig. 8D).

302 To examine this relationship across the population, we defined an afferent inputs similarity index for multiply innervated VS neurons, based on the phase of afferent 303 304 inputs and their EPSC amplitudes. The index ranges from 0-1, with smaller index 305 representing more divergent ESPC input tuning and larger index representing more similar tuning (see Methods). A classifier originally developed for visual cortical neurons 306 307 was used to quantify the tuning complexity of the postsynaptic neuron's membrane potential responses to sensory stimuli (Skottun et al., 1991). In this metric, neurons with 308 simple tuning show large AC and small DC responses, whereas complex cells exhibit 309 310 small AC and large DC responses (Fig. 8E). We found that the AC/DC ratio of the membrane potential was strongly correlated with the similarity index of afferent inputs 311 312 (Fig. 8F). In other words, convergence of more similarly tuned afferents yields a more 313 simple VS neuron response, and the convergence of more differently tuned afferents 314 generates a more complex postsynaptic response.

# 315 Spike tuning generation from similar and differently tuned afferents.

316 We next extended the comparison of presynaptic to postsynaptic tuning by 317 measuring the spiking responses of VS neurons during sensory stimulation. In a subset of VS neurons, the largest translational stimuli that we could deliver while holding the 318 319 cell was sufficient to evoke postsynaptic firing; in other neurons, a small bias current 320 was injected to evoke spiking during sensory stimulation (see Methods). Most VS neurons exhibited simple spike tuning, and received convergent inputs from similarly 321 322 tuned afferents (Fig. 9A). Some VS neurons with simple spike responses received convergent inputs from differently tuned afferents (Fig. 9B). Finally, complex spike 323 324 tuning in VS neurons was always generated by inputs from differently tuned afferents 325 (Fig. 9C). These three categories of input-output transformation (similar to simple, 326 different to simple, different to complex) were identified across all recordings from VS

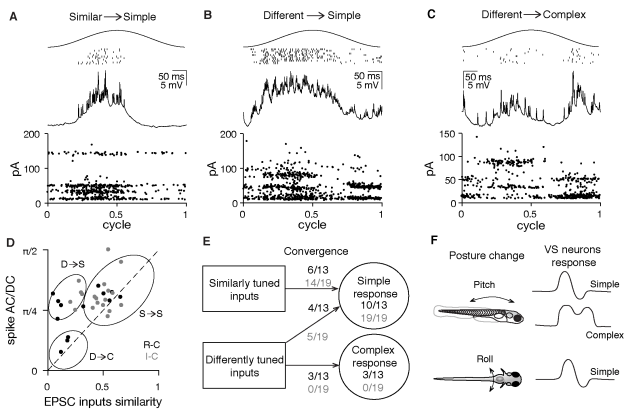


Figure 9: Spiking tuning complexity of VS neuron is partially explained by the similarity of tuning for the afferent inputs

A. Example cell showing that simple spiking tuning response is constructed from afferent inputs with similar tuning direction. Top, sensory-evoked spike raster of a VS neuron during 2 Hz, 0.02 g translational movement on the I(-)-C(+) axis (11 trials); middle, average membrane potential of the VS neuron (11 trials); bottom, sensory-evoked EPSC response (12 trials); each dot represents one EPSC.

B. As in A, for an example cell showing simple spiking tuning response arising from afferent inputs with different tuning directions on the I(-)-C(+) axis.

C. As in A, for an example cell with a complex spiking tuning response due to afferent inputs with different tuning directions on the R(+)-C(-) axis.

D. Correlation of EPSC input similarity index and spike activity AC/DC ratio, for all spiking VS neurons on the R-C axis (black, n=13 recordings) and I-C axis (gray, n=19). Circles, three categories of input-output transformation, corresponding to examples in A, B and C. Dashed, unity line. Pearson's R: 0.51

E. Quantification of input-output transformation of VS neurons; Ns represent individual cells.

F. Summary of different responses of VS neurons responding to posture change on the pitch and roll axes.

neurons (Fig. 9D). In total, most recordings (R-C, 6/13; I-C, 14/19) exhibited simple 327 328 spike tuning, of which 69% (20/29) received inputs from similarly tuned afferents, and 329 31% (9/29) received inputs from differently tuned afferents (Fig. 9E). Thus, convergence 330 of similarly tuned afferents yields simple spike tuning, but convergence of differently 331 tuned afferents can yield either simple or complex spike tuning. Interestingly, complex 332 spike tuning was only observed on the R-C axis (3/13 recordings), which subserves pitch movements, not the I-C axis (0/19), which subserves roll (Fig. 9F). These results 333 334 indicate that VS neurons may play different roles in maintaining body balance on

different body axes.

# 336 Discussion

# 337 Sensory convergence in the central vestibular nuclei

338 Taking advantage of the invariant synaptic transmission of electrical synapses, 339 we separated distinct afferent inputs that converge onto VS neurons and measured the 340 spatial and temporal tuning of each converging afferent in vivo. This analysis is 341 facilitated by the sparseness of connectivity, with < 6 afferents synapsing with each VS 342 neuron. These data resolve a conflict in the literature: anatomically, very few otolith 343 afferent terminals are observed in the lateral vestibular nucleus (Newlands and Perachio, 2003), but physiologically, afferent stimulation elicits monosynaptic EPSPs in 344 345 VS neurons (Boyle et al., 1992). Our data reveal that sparse but powerful afferent 346 synaptic contacts, located on the lateral dendrites of VS neurons, are sufficient to drive the membrane potential of the cell during sensory stimulation. Although these large 347 348 afferent inputs drive sensory responses, VS neurons receive a wealth of non-vestibular 349 synaptic contacts on their large dendritic arbors. This is consistent with previous findings that the activity of VS neurons is regulated by locomotion (Orlovsky, 1972), 350 proprioception (Neuhuber and Zenker, 1989), and other inputs (Sarkisian, 2000, Witts 351 352 and Murray, 2019). Interestingly, lateral geniculate neurons (LGN) of the visual 353 thalamus display a similar pattern of connectivity, with sparse, powerful afferent inputs 354 from retinal ganglion cells and weaker, diverse inputs from other sources (Usrey et al., 355 1999, Sherman, 2005). Our findings suggest that this configuration is common to VS 356 neurons as well.

357 Similarly tuned otolith afferents preferentially converge onto VS neurons (Fig. 7), 358 demonstrating that feedforward excitation can generate central neurons with simple 359 response properties. In a similar vein, thalamocortical inputs with similar angular tuning 360 also preferentially project onto the same site in somatosensory cortex, and the preferred tuning direction of the cortical neuron can be predicted by that of the presynaptic 361 362 thalamic neuron (Bruno et al., 2003). Furthermore, we found that convergence of differently tuned afferents can yield a more complex postsynaptic response in central 363 vestibular neurons, similar to bidirectional or complex tuning observed previously in cats 364 (Peterson, 1970) and primates (Angelaki and Dickman, 2000). This result generally 365 supports the hypothesized model (Angelaki, 1992) that the tuning of central vestibular 366 367 neurons can be constructed from cosine tuned inputs with varying tuning properties. 368 However, we find that convergence of differently tuned afferents can also yield simple 369 tuning in VS neurons (Fig. 8B), suggesting other factors such as inhibition (Straka and

Dieringer, 1996) and thresholding (Priebe et al., 2004) might be involved. We found no 370 371 evidence for polysynaptic excitatory circuits during afferent stimulation (Fig. S9 A and 372 B), and modelling indicates that excitatory synaptic input is sufficient to predict 373 subthreshold membrane potential and tuning (Fig. S9 C-F). However, stronger stimuli 374 might elicit inhibition and other nonlinearities. Across brain regions, sensory tuning of 375 central neurons is constructed by a variety of mechanisms. These include afferent convergence pattern (Alonso and Martinez, 1998, Priebe and Ferster, 2012), local 376 377 excitatory or inhibitory modulation (Wilent and Contreras, 2005), and nonlinear dendritic computation (Lavzin et al., 2012). Our results demonstrate that sensory response of a 378 379 central neuron can be constructed from the afferent inputs in a direct feedforward 380 manner.

# 381 Otolith afferent tuning properties in the larval zebrafish

The derived spatial tuning profile of afferents in the larval zebrafish is similar to 382 the polarity of the hair cells in otolith macula, consistent with results in fish (Fay, 1984, 383 Platt, 1977) and primates (Fernandez et al., 1972). Notably, tuning to dorsal or ventral 384 385 acceleration was relatively weak for most afferents, presumably due to the horizontal 386 orientation of the utricular membrane in larval zebrafish inner ear. Afferents were 387 preferentially tuned to contralateral acceleration (ipsilateral tilt) in the roll axis, consistent with the dearth of ipsilaterally tuned hair cells in larval zebrafish (Haddon et al., 1999). 388 In species with more centrally located line of polarity reversal (Fernandez and Goldberg, 389 390 1976a, Tomko et al., 1981), we would predict more convergence of oppositely tuned 391 afferents, and correspondingly more complex response of VS neurons in the roll axis, 392 as seen in cats (Peterson, 1970).

393 A significant question in vestibular systems is whether central vestibular neurons receive selective projections from afferents with regular as opposed to irregular firing. 394 395 Both regular and irregular afferents are thought to converge on VS and vestibulo-ocular 396 reflex neurons in mammals, based on studies comparing recruitment thresholds of 397 afferent inputs (Boyle et al., 1992). Our data provide direct evidence that vestibular inputs to VS neurons exhibit classic characteristics of irregular afferents (Eatock and 398 399 Songer, 2011): high-pass tuning, low spontaneous firing rate, and high CV of firing (Fig. S6). It is unknown whether regular utricular afferents exist in the larval zebrafish. 400 401 Although regular utricular afferents were observed in guitarfish (Budelli and Macadar, 402 1979), they appear absent in toadfish (Maruska and Mensinger, 2015) and sleeper goby (Lu et al., 2004). Based on serial section EM, many afferents make no contacts with VS 403 404 neurons (Fig. 4C), leading us to conclude that either regular afferents have not yet developed or that they do not contact VS neurons in the larval zebrafish. 405

# 406 Linear and fast synaptic transmission via gap junctions

Our data reveal that electrical synapses mediate linear synaptic transmission
from otolith afferents to the VS neurons. In contrast, synaptic transmission at
retinotectal afferents in larval zebrafish is mediated solely by glutamate (Smear et al.,
2007), suggesting that electrical synapses in vestibular afferents are perhaps not simply
a feature of early larval development but play an important role in circuit computations,

potentially via their amplitude invariant transmission. Interestingly, mammalian 412 413 vestibular afferent synapses also exhibit amplitude invariant transmission in the medial 414 vestibular nucleus (Bagnall et al., 2008) and cerebellum (Arenz et al., 2008, Chabrol et 415 al., 2015), but via specialized glutamatergic terminals (Turecek et al., 2017, McElvain et 416 al., 2015), indicating that frequency-independent transmission is a hallmark of vestibular 417 signaling across vertebrates. Furthermore, mixed electrical and chemical synapses have been anatomically identified between vestibular afferents and VS neurons in both 418 419 adult fish (Korn et al., 1977) and rodents (Nagy et al., 2013), suggesting the mixed 420 synapse may be a conserved mechanism across species to implement fast, frequency-421 independent transmission in the lateral vestibular nucleus. The amplitude invariance of this connection allowed us to examine whether there was any relationship between an 422 afferent's sensory gain or firing rate and the synaptic amplitude it evokes in a VS 423 424 neuron. No correlation appeared in either of these measures (Fig. S8), indicating that at least within this population, synapse size is not "normalized" by firing rate. 425

#### 426 VS pathway underlying sensorimotor transformation

427 The VS pathway is important for posture control. Larval zebrafish swim at high frequencies up to 100 Hz (McLean and Fetcho, 2009) and are naturally unstable in 428 429 water (Ehrlich and Schoppik, 2017). Our study examined the response of VS neurons with translational stimuli in the range of 0.04-0.12 g and 0.5-8 Hz, head movement 430 parameters comparable to slow swimming (Voesenek et al., 2016) or small angle tilting 431 432 motion in the larval zebrafish. In the roll axis all VS neurons are tuned to ipsilateral tilt 433 (Fig. S10), consistent with data from calcium imaging (Migault et al., 2018, Favre-Bulle 434 et al., 2018), suggesting they might excite specific motor units in the spinal cord to 435 produce compensatory movements (Bagnall and McLean, 2014). On the pitch axis, VS neurons have more heterogeneous responses, including simple tuning to either rostral 436 or caudal acceleration (Fig. S10), as well as complex responses (Fig. 9E) to both 437 438 directions. Thus, when the animal is destabilized by excessive nose-up or nose-down 439 tilts, VS neurons might activate non-specific motor units, increasing the likelihood of 440 swim bouts to regain balance (Ehrlich and Schoppik, 2017).

441 The high-pass tuning and phase lead of otolith afferents innervating VS neurons will make larvae most sensitive to ongoing changes in tilt or acceleration, especially at 442 443 high frequency. These data are consistent with behavioral observations that larvae 444 become more likely to swim to correct their position in the pitch axis when angular 445 velocity (i.e., changing tilt) reaches a critical threshold (Ehrlich and Schoppik, 2017). 446 This compensatory postural adjustment, which relies on both trunk and fins, is absent in rock solo larvae (Ehrlich and Schoppik, 2019), in line with our results on the absence of 447 sensory tuning in those animals. Larval VS neurons receive similar amounts of inputs 448 from rostrally and caudally tuned afferents, suggesting both nose-up and nose-down 449 450 tilts are equally detected by the VS pathway. In contrast, the vestibulo-ocular pathway 451 shows an anatomical bias for representation of nose-up body tilt (Schoppik et al., 2017). 452 This indicates that different strategies might be involved to adjust body posture and eye position for pitch movements in larval zebrafish. 453

454 Moreover, it is important for animals to distinguish self-generated and external 455 vestibular signals. We described the direct excitatory inputs from vestibular afferents 456 onto the VS neurons during passive movements. How do self-generated motion signals

457 modulate the activities of VS neurons? Projections from Purkinje cells in the cerebellum

- 458 are thought to suppress sensory-evoked activity in VS neurons during voluntary self-
- 459 motion (Cullen, 2019). In the future, it would be interesting to use *in vivo* whole-cell
- 460 physiology to investigate how central vestibular neurons distinguish self-generated
- 461 movements from passive movements.

# 462 Methods:

- 463 Fish lines and husbandry:
- 464 *Tg(nefma:LRL:Gal4)* was established by injecting the construct containing hsp70
- 465 promoter (Kimura et al., 2014), and the insertion site was set at the upstream of the
- 466 nefma gene with the CRISPR target sequence: CATCGACGGATCAATGG. The
- 467 Tg(nefma:gal4) fish line was generated by crossing Tg(nefma:LRL:Gal4) with a
- 468 ubiquitous-Cre fish. The otoge. 1522+2T>A -/- (rock solo) mutant is vestibular deficient
- due to a splice site mutation in *otogelin* (Mo et al., 2010, Roberts et al., 2017). Rock
- solo homozygotes on a *Tg(nefma:gal4, UAS:GFP)* background were crossed to rock
- 471 solo WT/heterozygotes to produce clutches containing WT, heterozygotes and
- 472 homozygotes for recording purposes. The rock solo homozygotes were identified by the
- absence of anterior otolith (utricle). All experiments and procedures were approved by
- the Animal Studies Committee at Washington University and adhere to NIH guidelines.
- Animals were raised and maintained in the Washington University Zebrafish Facility at
- 476 28.5°C with a 14:10 light:dark cycle. Larval zebrafish (4-7 dpf) were housed either in
- 477 petri dishes or shallow tank with system water. Adult animals were maintained up to 1
- 478 year old with standard procedure.
- 479 <u>Electrophysiology:</u>
- 480 VS neurons were targeted for whole-cell patch clamp recordings based on their
- 481 characteristic position and fluorescence in the *Tg(nefma:gal4, UAS:GFP)* fish. The
- 482 larvae (4-7 dpf) were paralyzed by 0.1%  $\alpha$ -bungarotoxin and embedded in a 10 mm
- 483 FluoroDish (WPI) with low-melting point agarose (Camplex SeaPlaque Agarose, 2.4%
- 484 in system water). Fish were immersed in extracellular solution ([in mM] NaCl 134, KCl
- 2.9, MgCl<sub>2</sub> 1.2, HEPES 10, glucose 10, CaCl<sub>2</sub> 2.1, osmolarity ~295 mOsm and pH ~
- 486 7.5) and a small piece of skin above the brainstem was carefully removed by sharpened
- tungsten pins. The fish was transferred to an epifluorescence microscope equipped with
   immersion objectives (Olympus, 40x, 0.8 NA), infrared differential interference contrast
- 489 optics and air-bearing sled recording table.
- 490 Patch pipettes (7-9 MΩ) were filled with internal solution ([in mM] K gluconate 125,
- 491 MgCl<sub>2</sub> 2.5, HEPES 10, EGTA 10, Na<sub>2</sub>ATP 4, Alexa Fluor 568 or 647 hydrazide 0.05-0.1,
- 492 osmolarity ~295 mOsm and pH ~ 7.5). After whole-cell configuration was achieved,
- 493 voltage clamp and current clamp signals were recorded at room temperature with a
- 494 Multiclamp 700B, filtered at 10 kHz (current clamp) or 2 kHz (voltage clamp), digitized at

50 kHz with Digidata 1440 (Axon Instruments), and acquired by Clampex 10 (MolecularDevices).

497 Before the vestibular stimulus was delivered to the fish, the immersion objective was

498 removed from the recording chamber. During the recording, series resistance was

499 monitored every 15 s to ensure good recording quality; neurons with series resistance

variation > 25% were discarded. After recording, the recorded cell was imaged with

501 epifluorescence to confirm cell identity.

# 502 <u>Vestibular stimulation:</u>

503 The recording rig was custom-designed to allow delivering user-controlled movement to

the fish during recording without losing whole-cell access. The microscope and a one-

axis or two-axis air-bearing sled (Aerotech, ABL1500WB or ABL1500&1500WB) were

506 fixed on an air table. Manipulators (Microstar) and recording platform (ThorLab) were

507 positioned on the sled. The sled was powered with the Aerotech transformers (TM5),

508 NPdrivers (NDRIVECP10-MXU&NDRIVECP20-MXU) and nitrogen gas (Airgas, NI

509 UHP300). Stimuli were designed in Matlab and imported into the program by Aerotech

510 software (Motion Designer/Composer), with additional tuning as required to compensate

511 for the motion of the underlying air table. Movement was recorded by an accelerometer

512 (Sparkfun, ADXL335) attached to the platform. Motion signals were digitized at 50 kHz

with Digidata 1440 (Axon Instruments), and acquired in Clampex 10 (Molecular

514 Devices).

515 Fish were embedded either dorsal side up (movements on rostral-caudal and ipsilateral-

516 contralateral axes) or left/right side up (movements on dorsal-ventral and rostral-caudal

axes) and a series of frequency-varying sinusoidal translational stimulus was applied.

The stimulus amplitude was set at 0.02 g or 0.06 g (min to max: 0.04 g or 0.12 g

respectively), and stimulus frequency range was 0.5-8 Hz. For spatial tuning

520 measurements, linear translation was applied on four different axes (0-180°, 45°-225°,

521 90°-270°, 135°-315°) on the horizontal plane. To record spike tuning in neurons without

522 spontaneous firing, a rheobase current was injected to depolarize the cell.

523 <u>Vestibular afferent stimulation and pharmacology</u>

A glass pipette electrode (2-5 MΩ) filled with extracellular solution ([in mM] NaCl 134,

525 KCl 2.9, MgCl\_2 1.2, HEPES 10, glucose 10, CaCl\_2 2.1, osmolarity ~295 mOsm and pH ~

526 7.5) was connected to a stimulator (A-M systems, Model 2100), and placed in the

vestibular ganglion to stimulate the vestibular afferents. A train of 0.1 ms, 1  $\mu$ A - 1 mA

528 electrical pulses at varying frequencies were delivered to elicit EPSCs in the recorded

529 cells. At least 20 trials of evoked EPSCs were recorded to establish a stable baseline.

530 AMPA receptors and gap junctions were blocked with 10  $\mu$ M NBQX and 500  $\mu$ M

- 531 carbenoxolone, respectively.
- 532 <u>Electron Microscopy:</u>

533 Ultrathin serial sections of brainstem from a 5.5 dpf zebrafish were a generous loan

from J. Lichtman and F. Engert. Using the published 18.8 nm/px reference map and

reconstructions (Hildebrand et al., 2017), we targeted re-imaging at 4 nm/px to the

entirety of the myelinated utricular afferents (identified by their peripheral processes 536

- 537 reaching for the utricular macula) and VS neurons (identified by their position and
- 538 axonal projections into the spinal cord) on one side of the brainstem, covering ~95  $\mu$ m
- 539 in the rostrocaudal axis. Imaging was carried out on a Zeiss Merlin 540 FE-SEM with a
- 540 solid-state backscatter detector. The ATLAS scan engine was controlled via WaferMapper (Hayworth et al., 2014). The resulting images were aligned onto the 18.8 541
- nm/px dataset using linear affine transformations in FIJI with the TrakEM2 plug-in 542
- 543 (Cardona et al., 2012) and will be freely available after publication. In a small subset of
- 544 identified synapses, we carried out further re-imaging at 1nm/px to visualize the
- 545 hallmarks of gap junctions.
- 546 The existing tracings of VS neurons and utricular afferents were extended to cover
- 547 branches that had been missed or untraced in the original dataset. Afferent/VS neuron
- appositions were considered to be synaptic contacts if the presynapse contained 548
- 549 vesicles, the membranes were tightly apposed and straight, and there were signs of a
- 550 postsynaptic density. In cases where appositions were more difficult to determine, such
- 551 as those parallel to the plane of section, vesicle clustering at a tight apposition was used
- 552 as the criterion for a synapse.
- 553 Analysis:
- 554 All analysis are implemented in Matlab (Mathworks).
- 555 Event detection:
- 556 EPSC events were detected by a derivative method (Bagnall and McLean, 2014).
- 557 Tuning index of all EPSCs was calculated as the vectoral sum of all events' phase,
- weighted by the EPSC amplitude. 558
- Tuning index =  $\frac{|\sum A_j * e^{i*\theta_j}|}{\sum A_i}$ ,  $i = \sqrt{-1}$ 559
- $(A_i \text{ is the amplitude of each EPSC event } j, \text{ and } \theta_i \text{ is the phase of that event relative to}$ 560
- the sinusoidal stimulus on each axis.) 561
- 562 Deconvolution of electrical and chemical signals

563 We assumed that the signals we observed on voltage clamp were majorly composed of electrical EPSCs and chemical EPSCs from afferents, based on our observation from 564 the pharmacology data. 565

- 566
- $S = conv(w_e, s_e) + conv(w_c, s_c) + residuals$ 567
- 568
- $(w_e, w_c \text{ are the kernels of electrical and chemical components of EPSCs, both derived$ 569
- from their waveforms shown in Fig. 2, and  $s_e$ ,  $s_c$  are the separated electrical and 570
- chemical signals) 571

572 A sparse deconvolution algorithm with L1 regularization, derived from FISTA ((Beck and 573 Teboulle, 2009), was applied to obtain  $s_e$ ,  $s_c$  by minimizing the objective function:

$$F(s_e, s_c) \equiv \frac{1}{2} (S_{observed} - S_{reconstruct})^2 + \lambda_e * |s_e| + \lambda_c * |s_c|,$$

575 where:

$$S_{reconstruct} = conv(w_e, s_e) + conv(w_c, s_c)$$

576 577

578  $\lambda_e$  and  $\lambda_c$  were defined by the root mean square of the signal and magnitude of kernel 579 waveform:  $\lambda_i = rms(S_{observed}) * \sqrt[2]{w_i * w_i^T}$ . Maximum iteration cycle was set at 500

# 580 <u>Clustering and quantification of electrical events</u>

581 Amplitude-invariant EPSCs are mediated by gap junctions, therefore only electrical

signals  $s_e$  were used to infer individual afferent inputs. A threshold of  $3.5 * std(s_e)$  was

used for event detection. Detected electrical events were clustered by ISO-SPLIT

584 (Chung et al., 2017). Some clusters were merged or split manually after examination.

585 Clusters with refractory period (threshold: probability < 0.003 within 1 ms) in auto-

586 correlograms (100 ms) were considered from an individual afferent.

587 For each cluster, the tuning vector of inferred afferent k on each axis was quantified as:

588 
$$z_k = \frac{\sum e^{i*\theta_j}}{N_c * A_s} * f_s, i = \sqrt{-1}$$

589 ( $\theta_j$  is the phase of EPSC event *j* in cluster *k*.  $N_c$  is the number of cycles for sinusoidal 590 translation,  $f_s$  [Hz] and  $A_s$  [g] are the frequency and amplitude of the stimulus. The 591 absolute value and argument of *z* represent the tuning gain and the tuning phase, 592 respectively.)

593 Tuning in four axes was fitted into a 2-dimensional spatiotemporal model (Angelaki, 594 1992) to obtain the maximum tuning direction, the tuning gain and phase in that 595 direction.

596 Afferent inputs similarity index for a VS neuron was determined as:

597 
$$\frac{|\sum A_k * z_k|}{\sum A_k * n_k}$$

598 ( $A_k, z_k, n_k$  are the average EPSC amplitude, tuning vector and number of events for 599 cluster *k*.)

# 600 AC/DC response quantification

AC of membrane potential and spiking response were defined as the amplitude of

sinusoidal fit (2 Hz) of the membrane potential, and the spike vectorial sum during

603 sensory stimulation, respectively. DC of membrane potential and spiking response were

604 defined as the average membrane potential during sensory stimulation above baseline

605 (no stimulation), and the total spike number during sensory stimulation above baseline.

606 For spiking responses, VS neurons with firing rate > 4 spike/cycle and spike AC or

- 607 DC >1 spike/cycle were included in the analysis.
- 608 <u>Bootstrapping</u>:  $\binom{n_i}{2}$  afferent pairs were counted for VS neuron *i* with  $n_i$  distinct afferent
- inputs  $(n_i \ge 2)$ . The same number of total afferent pairs  $\sum_{i=1}^{m} {n_i \choose 2}$  from all *m* VS neurons
- 610 was randomly selected among all  $\sum_{i=1}^{m} n_i$  inferred afferents to determine the
- 611 convergence angle or phase difference distribution by chance, and such selection was
- 612 performed 5000 times to calculate mean and standard deviation.

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- 624

# 625 Author contributions:

- 626 K.M and S.H generated the *Tg(nefma:gal4, UAS:GFP)* fish line. Z.L and M.B conceived
- the project. Z.L performed the electrophysiology, confocal imaging experiments and
- analyzed the data. T.H helped develop the deconvolution algorithm for sorting EPSC
- events. M.B, J.M and D.H carried out the serial section EM imaging and reconstruction.
- 630 Z.L. and M.B wrote the manuscript with input from all other authors.
- 631

# 632 **Declaration of Interests**:

- 633 The authors declare no competing interests
- 634

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