

Motor neurons are dispensable for the assembly of a sensorimotor circuit for gaze stabilization

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1 **Sensorimotor reflex circuits engage distinct neuronal subtypes, defined by precise connectivity, to transform sensation into**
2 **compensatory behavior. Whether and how motor neuron populations specify the subtype fate and/or sensory connectivity**
3 **of their pre-motor partners remains controversial. Here, we discovered that motor neurons are dispensable for proper con-**
4 **nectivity in the vestibular reflex circuit that stabilizes gaze. We first measured activity following vestibular sensation in pre-**
5 **motor projection neurons after constitutive loss of their extraocular motor neuron partners. We observed normal responses**
6 **and topography indicative of unchanged functional connectivity between sensory neurons and projection neurons. Next,**
7 **we show that projection neurons remain anatomically and molecularly poised to connect appropriately with their down-**
8 **stream partners. Lastly, we show that the transcriptional signatures that typify projection neurons develop independently**
9 **of motor partners. Our findings comprehensively overturn a long-standing model: that connectivity in the circuit for gaze**
10 **stabilization is retrogradely determined by motor partner-derived signals. By defining the contribution of motor neurons**
11 **to specification of an archetypal sensorimotor circuit, our work speaks to comparable processes in the spinal cord and ad-**
12 **vances our understanding of general principles of neural development.**

13 INTRODUCTION

14 Developing sensorimotor reflex circuits must precisely connect functional subtypes of neurons to ensure appropriate behavior.
15 For example, withdrawal from noxious stimuli requires maturation of a sensorimotor circuit that uses subtypes of spinal interneu-
16 rons to transform noxious stimulation into activation of both ipsilateral flexor and contralateral extensor motor neurons¹. Work
17 over the past 40 years has highlighted motor partner populations as possible orchestrators of connectivity in pre-motor reflex cir-
18 cuits²⁻⁶, but controversy remains about the nature of their role. In the spinal cord, molecular perturbations of motor neuron iden-
19 tity have provided evidence both for⁷⁻¹¹ and against¹²⁻¹⁶ an instructive role in determining pre-motor fate. Part of this contro-
20 versy stems from the wide variety of inputs to spinal motor neurons¹⁷, the molecular and functional heterogeneity of pre-motor in-
21 terneurons^{14,15}, and their complex roles in gait and posture¹⁸. Further, transcription factors play multivariate and redundant roles
22 in spinal motor neuron development^{19,20}, such that the effects of molecular perturbations of identity can be masked.

23 The sensorimotor circuit for vertical gaze stabilization offers a simple framework to evaluate whether and how motor neurons shape
24 pre-motor circuit fate and connectivity. The vertebrate vestibulo-ocular reflex circuit consists of three neuron types – peripheral
25 sensory, central projection, and extraocular motor neurons – that stabilize gaze after head/body tilts (Figure 1A)²¹. Subtype fate,
26 anatomical connectivity, and function are inextricably linked: directionally-tuned sensory neurons innervate nose-up/nose-down
27 subtypes of projection neurons, which in turn innervate specific motor neurons that selectively control either eyes-down or eyes-up
28 muscles^{3,22-29}. As both the recipients and origin of directional information, projection neuron fate specification is tantamount to
29 proper circuit assembly. Recent work has established the vertical vestibulo-ocular reflex circuit in zebrafish as a model to uncover
30 determinants of fate and connectivity^{24-26,29} given the ease of optical imaging, abundant tools for genetic perturbations, rapid de-
31 velopment, and robust evolutionary conservation.

32 The current model for vestibulo-ocular reflex circuit development was motivated by pioneering work in chick^{3,30} and formalized by
33 Hans Straka: “[circuit assembly] is accomplished by a specification process that retrogradely transmits post-synaptic target identi-
34 ties to pre-synaptic neurons.”³¹. In its strongest form, this “retrograde” model posits a causal role for extraocular motor neurons in
35 specifying the fate (sensory selectivity) of central projection neurons. This key prediction – that loss of motor neurons would disrupt

36 sensory selectivity in pre-synaptic projection neurons – remains untested. In zebrafish, extraocular motor neurons are temporally
37 poised for such a role. Motor neurons are organized into spatial pools, and though synaptogenesis at ocular muscle targets begins
38 late in development³², motor neuron fate (muscle target and pool location) is determined early³³. Projection neurons are born at
39 roughly the same time as motor neurons and extend axons shortly afterwards, poising them to receive deterministic signals that
40 could retrogradely specify their sensory selectivity²⁹.

41 Here, we adopted a loss-of-function approach to determine whether motor partner populations specify identity or instruct con-
42 nectivity across an entire vestibular reflex circuit in zebrafish. We generated a new mutant allele for the *phox2a* gene to eliminate
43 the extraocular motor neurons used for vertical gaze stabilization. Combining functional, anatomical, and sequencing approaches,
44 we then demonstrated that motor neurons are dispensable for three aspects of pre-motor reflex circuit assembly: (i) directionally-
45 appropriate connectivity between sensory and projection neurons, (ii) assembly of projection neurons with motor partners, and
46 (iii) the transcriptional profiles of projection neurons. The current model of vestibulo-ocular reflex circuit development must there-
47 fore be revised: up/down projection neuron subtype fate cannot be retrogradely established by a motor partner-derived signal. In-
48 stead, the signals that specify fate must lie elsewhere. More broadly, our work argues against a deterministic role of motor neurons
49 in specifying the fate and sensory connectivity of pre-motor circuit components.

50 RESULTS

51 **Constitutive loss of *phox2a* prevents extraocular motor neuron specification and impairs vertical gaze stabilization behavior**

52 Extraocular motor neurons for vertical/torsional gaze stabilization are located in cranial nuclei III (nIII) and IV (nIV). To eliminate
53 nIII/nIV motor neurons and by extension, any secreted signals, we used a genetic loss-of-function approach (Figure 1A). A single
54 highly-conserved transcription factor, *phoxa*, specifies nIII/nIV fate^{34–37}. In the vestibulo-ocular reflex circuit, *phox2a* is exclusively
55 expressed in nIII/nIV motor neurons but not its upstream partners (Figure 1B). Therefore, *phox2a* is an ideal genetic target to elimi-
56 nate motor-derived signals without compromising evaluations of upstream functional development.

57 Prior mutagenesis established a *phox2a* loss-of-function allele in zebrafish³⁴, but the line has since been lost. Here, we generated
58 three new *phox2a* loss-of-function alleles using CRISPR/Cas9 mutagenesis (Figure 1C) (one allele shown here; additional alleles de-
59 scribed in Methods). Consistent with prior reports and human mutations³⁸, both eyes in *phox2a* null mutants were exotropic (ro-
60 tated towards the ears) reflecting a loss of motor neurons in nIII/nIV. *phox2a* mutants failed to hatch from their chorions without
61 manual intervention and did not inflate their swim bladders by 5 days post-fertilization (dpf) (Figure 1D), phenotypes not previously
62 reported³⁴. Consequently, null mutants do not survive past 7 dpf. We did not observe these morphological phenotypes in wildtype
63 and heterozygous siblings (Figure 1D). As vestibulo-ocular reflex circuit architecture and behavior is established by 5 dpf^{24,25,29},
64 premature lethality did not preclude further measurements of circuit development.

65 To validate *phox2a* loss-of-function, we leveraged a downstream transcription factor: *isl1*³⁹. The *Tg(isl1:GFP)* line⁴⁰ labels all nIII/nIV
66 motor neurons except inferior oblique neurons³³, which comprise one of four pools for upwards eye rotations. We first quanti-
67 fied changes in the number of labeled nIII/nIV neurons (Figure 1E-Figure 1F). In *phox2a* mutants, we observed an expected and
68 near-total loss of *isl1* expression (WT: 298±19 neurons across both hemispheres; null: 19±11 neurons; Wilcoxon rank sum test,
69 $p=2.5\times 10^{-4}$) at 5 dpf, well-after nIII/nIV differentiation is complete³³. Unexpectedly, we also observed slightly fewer neurons in
70 *phox2a* heterozygotes (heterozygote: 229±20 neurons; Wilcoxon rank sum test against WT, $p=6.7\times 10^{-4}$). In heterozygotes, loss
71 of *isl1* fluorescence was restricted to the medial domain of dorsal nIII, which contains some of the earliest-born neurons in nIII/nIV
72 (Figure S1A-Figure S1C)³³. Globally, this manifested as a rostral and ventral shift in the positions of all neurons mapped (Fig-
73 ure S1D) (two-sample, two-tailed KS test, WT vs. heterozygotes: mediolateral axis, $p=0.13$; rostrocaudal: $p=4.0\times 10^{-29}$; dorsoventral:
74 $p=2.5\times 10^{-9}$). This region contains two motor pools that control the inferior (IR) and medial rectus (MR) muscles³³. We conclude
75 that *phox2a* acts in a dose- and birthdate-dependent manner to specify nIII motor pool fate.

76 Together, these observations validate our *phox2a* loss of function alleles as a selective means to disrupt nIII/nIV motor neuron fate
77 specification and vertical eye rotation behavior.

78 **Peripheral-to-central circuit assembly does not require motor partners**

79 Vertical gaze stabilization requires that (1) peripheral VIIIth nerve sensory afferents relay tilt sensation (nose-up/nose-down) di-
80 rectly to projection neurons in the tangential nucleus, and (2) projection neurons innervate appropriate nIII/nIV pools (eyes-up/eyes-
81 down). For proper circuit function, appropriate connectivity must first develop across up/down circuit subtypes. The “retrograde”
82 model predicts that motor partners specify circuit assembly. Therefore, in the absence of motor neurons, projection neurons should
83 fail to respond selectively to directional tilt sensations – either due to loss of their fate, the fate of upstream sensory afferents, or
84 sensory-to-central connectivity.

85 To evaluate upstream circuit formation, we measured tilt-evoked responses in projection neurons using Tilt-In-Place Microscopy
86 (TIPM)⁴¹ (Figure 2A-Figure 2B). Peripherally, tilts activate utricular VIIIth nerve sensory inputs to projection neurons^{29,41}. We used
87 a galvanometer to deliver tonic nose-up and nose-down pitch tilts to *phox2a* null larvae and sibling controls. We then measured
88 the activity of a calcium indicator, GCaMP6s⁴², in projection neurons. We performed experiments at 5 dpf, when nearly all projec-
89 tion neurons are selective for one tilt direction²⁹, circuit architecture is stable²⁵, and gaze stabilization behavior is directionally-
90 appropriate²⁴.

91 Projection neuron responses and topography were strikingly unchanged in *phox2a* mutants compared to controls. We recorded
92 the activity of n=297 neurons from N=16 *phox2a* mutants and n=440 neurons from N=21 sibling controls (Methods and Table 1
93 split by genotype). We observed comparable ratios of projection neuron subtypes (sib: 46% nose-down, 47% nose-up, 7% untuned;
94 *phox2a*: 49% nose-down, 44% nose-up, 7% untuned) (Figure 2C). Next, we evaluated their topography (Figure 2D-Figure 2E). Pro-
95 jection neurons are topographically organized along the dorso-ventral axis by their directional selectivity²⁹. Global spatial separa-
96 tion between subtypes remained significant in *phox2a* mutants (one-way multivariate analysis of variance, p=0.004). We also com-
97 pared the topography of nose-up and nose-down neurons separately across *phox2a* genotypes. Nose-down neurons were com-
98 parably distributed between null and control larvae (one-way multivariate analysis of variance, p=0.15). We observed a minor lat-
99 eral shift to nose-up neurons in null mutants (median mediolateral position, sib: 15.2μm from medial edge; *phox2a*, 13.2μm; two-
100 tailed, two-sample KS test, p=3.0x10⁻⁴) but no changes in other spatial axes (dorsoventral: p=0.16; rostrocaudal: p=0.56). The small
101 medial deviation (2μm across a 40μm space) is within the limits of our registration error. We conclude that projection neuron to-
102 pography is established independently of motor partners.

103 Projection neuron sensitivity and selectivity also developed comparably between *phox2a* mutants and siblings (Figure 2F-
104 Figure 2K). Projection neurons responded to tilt sensations with comparable magnitudes (Figure 2G-Figure 2J) (nose-down mean
105 ΔFF, sib: 1.86±1.69; *phox2a*: 2.07±1.48; two-tailed Wilcoxon rank sum test, p=0.98; nose-up mean ΔFF, sib: 1.24±1.23; *phox2a*:
106 1.02±0.89; p=0.18). Previously, we defined a metric to describe a neuron’s selectivity for one tilt direction (0 = equal responses
107 to up/down; 1 = maximally selective)²⁹. Directional selectivity remained unchanged in *phox2a* mutants (Figure 2H-Figure 2K)
108 (nose-down mean index, sib: 0.73±0.29; *phox2a*: 0.68±0.29; two-tailed Wilcoxon rank sum test, p=0.85; nose-up mean index, sib:
109 0.85±0.26; *phox2a*: 0.81±0.29; p=0.12). Collectively, this demonstrates that the functional responses of projection neurons and, by
110 inference, connectivity with utricular afferents are not shaped by motor partners.

111 Ventral projection neurons receive additional input from the semicircular canals²⁹, which encode phasic (fast) tilt sensation. To
112 activate sensory afferents from the semicircular canals, we used TIPM to deliver two impulses of angular rotation (Figure 3A-
113 Figure 3B)^{29,41}. We observed no changes in *phox2a* mutants. Projection neurons responded to impulses in comparable ratios (Fig-
114 ure 3E) (sib: 58% responsive; *phox2a*: 71% responsive). Responsive projection neurons remained localized to the ventral nucleus
115 (dorsoventral axis: two-tailed, two-sample KS test, p=0.99). Lastly, the functional properties of projection neurons were unchanged
116 (Figure 3F-Figure 3G). We observed no change in calcium response magnitudes (Figure 3F) (mean ΔFF, sib: 0.33±0.29; *phox2a*:
117 0.36±0.40; two-tailed Wilcoxon rank sum test, p=0.85) or lack of directional selectivity (Figure 3G) (mean index, sib: 0.07±0.41;
118 *phox2a*: 0.06±0.40; p=0.39). Therefore, fate and connectivity between phasic sensory afferents and projection neurons must not
119 require motor partners.

120 Lastly, we considered whether loss of one subtype of nIII/nIV neurons might alter connectivity. For example, loss of eyes-down mo-

121 tor pools could impair wiring between their corresponding nose-up sensory and projection neuron partners. Here, we leveraged
122 *phox2a* heterozygotes, which lack a subtype of nIII neurons (IR/MR) that contribute to downwards eye rotations (Figure S1). We ob-
123 served no differences in tonic tilt responses between *phox2a* wildtype, heterozygote, and null larvae, though we did note a minor
124 decrease in response strength to impulses (statistics in Table 1). We note that *phox2a* heterozygotes do not lack all motor pools for
125 downwards eye rotations. Nevertheless, we conclude that individual motor pools do not meaningfully contribute to connectivity be-
126 tween sensory and projection neurons.

127 Taken together, these experiments demonstrate intact directional selectivity for two peripheral sensory inputs – utricular and semi-
128 circular canal VIIIth nerve afferents – and appropriate connectivity with projection neurons. We conclude that functional sensory-to-
129 central circuit formation is established independently of motor partners.

130 **Projection neurons remain competent to assemble with appropriate motor targets**

131 Motor partners could secrete signals that initiate pre-motor axon outgrowth, target arriving axons to specific motor pools, or trigger
132 synaptogenesis³. Motor pool topography in nIII/nIV reflects ocular muscle targets: dorsal pools innervate downward-rotating mus-
133 cles (superior oblique and inferior rectus), while ventral pools target the converse (eyes-up, superior rectus and inferior oblique)^{33,43}.
134 In turn, projection neuron somatic and axonal organization mirrors motor pool topography^{28,29}, which could facilitate directionally-
135 selective circuit assembly. We reasoned that projection neurons may fail to initiate axon outgrowth, target spatially-appropriate mo-
136 tor pools, and/or form synapses in *phox2a* mutants. To test this hypothesis, we measured changes in projection neuron anatomy at
137 5 dpf, when axonal arbors are established and stable²⁵.

138 To test whether projection neurons establish gross, long-range (hindbrain to midbrain) axonal outgrowths, we performed optical
139 retrograde labeling⁴⁴ using a photolabile protein, Kaede. We targeted the medial longitudinal fasciculus at the midbrain-hindbrain
140 boundary, which contains projection neuron axons^{25,29} (Figure 4A). In both *phox2a* mutants and sibling controls, we observed
141 retrograde photolabeling of projection neuron soma (Figure 4B), supporting that initial axon outgrowth does not require motor
142 partner-derived signals.

143 Next, we evaluated whether projection neuron axons remain capable of wiring with spatially-appropriate motor partners. Projec-
144 tion neuron axons segregate along the dorsal (nose-up) and ventral (nose-down) axes according to their birth order (early/late born,
145 respectively)^{28,29} and the pool topography of their motor targets³³. To test whether projection neurons retain this topography,
146 we optically labeled the axons of early-born (before 30 hpf) projection neurons²⁹. In *phox2a* mutants, axons remained dorsoven-
147 trally segregated at midbrain targets (Figure 4C, inset). Typically, projection neurons robustly collateralize to nIII/nIV targets at the
148 midbrain-hindbrain boundary. We did not observe collaterals to nIII/nIV in *phox2a* mutants (Figure 4C). However, projection neu-
149 rons still robustly arborized to more rostral, spinal-projecting targets in the nucleus of the medial longitudinal fasciculus, suggesting
150 they retain the machinery necessary to collateralize. Consistent with this hypothesis, we observed that projection neurons formed
151 occasional, small collaterals in *phox2a* mutants with few (1-5%) nIII/nIV neurons remaining (Figure 4D). We conclude that projec-
152 tion neurons remain competent to assemble with spatially-appropriate targets.

153 If motor neurons are required to initiate synaptogenesis, then projection neurons should fail to develop pre-synaptic machinery.
154 To test this hypothesis, we performed fluorescent *in situ* hybridization against common pre-synaptic transcripts: synaptophysin a
155 (*syph*), synaptic vesicle glycoprotein (*sv2*), and synapsin I (*syn1*). In both *phox2a* mutants and controls, we observed robust tran-
156 script expression in projection neuron somata at 5 dpf (Figure 4E), well-after synaptogenesis onset in wildtype larvae²⁹. Motor
157 partner-derived signals are thus not required for projection neurons to develop the necessary components for synaptogenesis.

158 Though motor neurons may play later roles in selecting and/or refining pre-motor input specificity, our data supports that projec-
159 tion neurons remain anatomically and molecularly poised to assemble with appropriate targets. We predict that absent collaterals
160 and synapses reflect a lack of adhesive contact necessary to stabilize^{45,46}, but not instruct the formation of nascent structures.

161 The transcriptional profiles of projection neurons are intact in the absence of motor partners

162 Functional and anatomical connectivity, from peripheral sensors to motor targets, develop independently of motor partners. Fate
163 in the vestibulo-ocular reflex circuit follows from connectivity²¹, but neuronal fate can also be defined with respect to unique tran-
164 scriptional signatures. Previously, we developed a sequencing pipeline to discover transcription factors that specify functional sub-
165 types of spinal motor neurons and evaluate the consequences of perturbations on transcriptional fate⁴⁷. We adapted this approach
166 to determine if loss of motor-derived signals changed the transcriptional profiles of projection neurons.

167 We compared the transcriptional profiles of projection neurons in *phox2a* mutants and sibling controls (Figure 5A) using bulk
168 RNA sequencing. We performed sequencing experiments at 72 hours post-fertilization (hpf), after projection neuron differ-
169 entiation is complete and synaptogenesis to motor targets has peaked²⁹. We sequenced projection neurons labeled by *Tg(-*
170 *6.7Tru.Hcctr2:GAL4-VP16);Tg(UAS-E1b:Kaede)*^{24,25,48,49} (Methods, Figure 5B, Figure S2). Neurons labeled in this line include, but
171 are not exclusive to the projection neurons in the tangential nucleus used for vertical gaze stabilization. Therefore, we evaluated our
172 bulk RNA sequencing dataset in the context of a single-cell reference atlas derived from the same transgenic line (Methods, Fig-
173 ure S3) to minimize noise from other labeled populations. We used *evx2*⁵⁰ as a reference, as it was expressed in all projection neu-
174 rons (Figure S3D) and highly detected (50%) in singly-profiled projection neurons.

175 There were strikingly few differentially-expressed genes in projection neurons between *phox2a* siblings and null mutants (Fig-
176 ure 5C, Table 2). All candidate differentially-expressed genes were lowly-expressed (detected in <10% of reference projection neu-
177 rons, Figure 5D). To determine if any candidates were differentially expressed in projection neurons, we used a fluorescent *in situ* hy-
178 bridization method⁵¹ in which fluorescence intensity correlates with detected transcript expression reliably across individual larvae
179 (Figure S4). We evaluated 8 candidate genes (Figure 5E-Figure 5F). Qualitatively, we observed no differences in expression patterns
180 between *phox2a* mutants and siblings, neither in candidates with significant differential expression in the bulk RNA sequencing
181 dataset or in a highly-expressed control markers, *evx2*.

182 We considered that our inability to detect differentially-expressed genes could arise from our exclusion of candidates based on their
183 expression in our reference single-cell atlas. Therefore, we repeated our analyses in unfiltered bulk sequencing data. The top 50
184 highest-expressed genes in *phox2a* siblings were highly detected in singly-profiled neurons labeled by *Tg(-6.7Tru.Hcctr2:GAL4-*
185 *VP16);Tg(UAS-E1b:Kaede)*, including projection neurons (Table 3). This suggests that our dissections adequately captured our tar-
186 get population. However, we again identified few differentially-expressed genes in our unfiltered data (Figure S5A-Figure S5C), with
187 substantial decreases as significance stringency increased. *In situ* hybridization validated that top candidates remained lowly ex-
188 pressed in projection neurons in both *phox2a* siblings and mutants (Figure S5D). Importantly, nearly all candidates had low de-
189 tection across all neurons in our reference single-cell atlas and had predicted expression in populations such as glia and the cau-
190 dal hindbrain (Methods, Table 4). Notably, some candidates were highly expressed in the medial vestibular nucleus, which lies on
191 the medial edge of the tangential nucleus and expressed *phox2a* (Figure S6). Together, we conclude that the any differential gene
192 expression in our data either reflects noise or contamination from other labeled populations, but not projection neurons in the tan-
193 gential nucleus.

194 We acknowledge the possibility that our *in situ* method is insufficiently quantitative to detect subtle differences in expression. Simi-
195 larly, despite using both bulk and single-cell RNA sequencing approaches, we may lack the resolution to uncover differential gene
196 expression within projection neurons. Nevertheless, consistent with functional and anatomical characterization, our sequencing
197 data argues that projection neurons acquire the correct transcriptional profiles in the absence of motor partner-derived signals.
198 Our findings are reminiscent of recent reports that the molecular signatures of spinal interneurons develop independently of mo-
199 tor partners¹⁵.

200 DISCUSSION

201 Here, we show that motor neurons are dispensable for fate specification in a canonical sensorimotor circuit. We first demonstrated
202 that peripheral sensory and central projection neurons develop appropriate, directionally-selective connectivity and topography in-

203 dependently of their motor partners. Next, we established that projection neurons remain anatomically and molecularly competent
204 to assemble with motor partners. Lastly, we show that loss of motor neurons does not meaningfully alter the transcriptional signa-
205 tures of their pre-motor projection neuron partners. By providing causal evidence against an instructional role of motor partners for
206 sensory connectivity, our work forces a revision of the current model for vestibulo-ocular reflex circuit formation. As proper connec-
207 tivity across multiple synapses is foundational for proper function, our work speaks to general mechanisms responsible for sensori-
208 motor circuit assembly.

209 **Transcriptional influences on motor neuron fate specification**

210 While the primary focus of our work was circuit assembly, we found that, unexpectedly, *phox2a* acts in a dose-dependent manner
211 to specify extraocular motor pool fate. Key evidence comes from *phox2a* heterozygotes, in which the earliest-born dorsal neurons
212 in nIII are lost but later-born neurons in nIII/nIV are intact. This observation extends prior characterizations of *phox2a* mutations in
213 zebrafish³⁴, chick³⁶, and human^{37,38}. Prior work hypothesized that *phox2a* dosage may regulate midbrain motor neuron differ-
214 entiation into visceral and somatic types³⁶. In other systems, transcription⁵²⁻⁵⁴, growth⁵⁵ and axon guidance factors⁵⁶ can act in
215 such a graded manner to regulate coarse cell type specification and wiring specificity. We extend these ideas to show that *phox2a*
216 dose-dependency acts both over closely-related subtypes (pools within a single cranial nucleus) and along a temporal axis, where
217 partial dosage preferentially targets the earliest-born neurons³³. Specifically, if *phox2a* is expressed in neural progenitor cells that
218 give rise to nIII/nIV, then the earliest-born motor neurons would have the shortest exposure to *phox2a*.

219 Molecular insight into ocular motor neuron pool specification is sparse but would be welcome given the strong links between ge-
220 netic development and ocular motor disease^{38,57-59}. For example, subpopulation markers could resolve the topography of pools
221 within dorsal nIII; whether IR/MR pools are spatially segregated or intermingled^{33,60}; whether the medial/lateral axis reflects func-
222 tional differences among motor neuron subtypes; and whether/how local interactions between motor neuron pools contributes to
223 fate specification⁶¹. In spinal circuits, the rich molecular understanding of motor pool specification^{47,62-65} has enabled targeted
224 perturbations of pool identity, allowing for major discoveries of their roles in circuit assembly^{9,13,15,66}. Our findings thus represent a
225 step forward towards understanding how developmental deficits may contribute to ocular motor disorders⁶⁷.

226 **Motor neurons: active or passive architects of pre-motor connectivity?**

227 Our discoveries advance outstanding controversies over whether motor neurons actively or passively shape pre-motor connectiv-
228 ity. We find that extraocular motor neuron axons do not serve as “pioneers”⁶⁸⁻⁷², with pre-motor axon targeting following passively
229 from motor-derived pathfinding signals². Such a model predicts that projection neuron targeting would be entirely ablated after
230 constitutive loss of extraocular motor neurons and their secreted signals⁷³⁻⁷⁷. Instead, we observed that projection neurons still es-
231 tablish long-range (hindbrain to midbrain) axonal projections, with appropriate spatial segregation that matches the topography
232 of their motor partners^{28,29,33}. Our findings complement reports in spinal circuits that pre-motor targeting is grossly appropriate
233 after manipulating the spatial source of, but not ablating, potential pathfinding signals¹³, and that the transcriptional fate of pre-
234 motor projection neurons similarly develops independently¹⁵. We point to the late development of ocular musculature^{78,79} com-
235 pared to spinal musculature² as a potential source of the dispensability of muscle-derived signals.

236 Our work is also inconsistent with the strongest form of the “retrograde” hypothesis for vestibulo-ocular reflex circuit assembly.
237 Originally, the retrograde model posited that motor neurons release a diffusible or cell-surface available signal that instructs pre-
238 motor collaterals to sprout and then innervate specific pools, enabling behavioral specificity^{3,31}. Here, the proper spatial and tem-
239 poral segregation of projection neuron axons suggests they remain poised to wire with spatially-appropriate (dorsal/ventral pools)
240 targets. Additional evidence comes from incomplete *phox2a* knockouts (1-5% of nIII/nIV remaining), where projection neurons
241 still form collaterals, though not robustly or reliably. We predict that projection neuron axons do not require a target-derived cue
242 to grow, search, and synapse onto motor targets, and simply lack the adhesive contact necessary to stabilize nascent structures (re-
243 viewed in^{45,46}).

244 Nevertheless, extraocular motor neurons might still play an active or passive role in selecting and/or refining input specificity from

245 their projection neuron partners. In spinal circuits, motor pool position passively imposes geometric constraints on pre-motor axon
246 targeting^{13,14}, and manipulating the dendritic structure of motor neuron axons transforms input specificity^{9,11}. Genetic perturba-
247 tions of nIII/nIV motor neuron position selectively compromise ocular responses to directional visual stimuli⁶¹, though the circuit-
248 level origin of such impairments is unclear. For the vestibulo-ocular reflex circuit, transforming all motor pools to the same fate or
249 genetically “scrambling” pool position could resolve whether motor input specificity is truly hard-wired in projection neurons, or
250 whether projection neurons instead target gross spatial domains irrespective of partner identity¹³. Motor neuron-derived signals
251 are of course capable of shaping their input by strengthening/weakening their inputs. Importantly, our results suggest that such
252 signals will not define the fate of projection neurons, and by extension, circuit architecture.

253 We note that our study does not eliminate one additional source of post-synaptic partner signals to projection neurons. As in pri-
254 mates⁸⁰, projection neurons also contact neurons in the interstitial nucleus of Cajal, also known as the nucleus of the medial longi-
255 tudinal fasciculus (INC/nMLF)²⁴. INC/nMLF neurons project early in development⁸¹ to spinal circuits used for postural stabilization
256 during swimming⁸²⁻⁸⁶. Notably, ablation of projection neurons disrupts postural stability⁵⁰. As we did not observe postural deficits
257 in *phox2a* mutants, we infer that projection neuron connectivity to INC/nMLF targets is present and functional. Correspondingly,
258 the development of projection neuron collaterals and synapses to INC/nMLF neurons appeared qualitatively normal in *phox2a* mu-
259 tants, supporting our interpretation that projection neurons retain the capacity to properly assemble with post-synaptic targets
260 even though similar structures to extraocular motor neurons are absent. In the future, if a similarly specific marker like *phox2a* is
261 identified that labels the INC/nMLF, it will be possible to test whether these neurons play a role in vestibulo-ocular reflex circuit de-
262 velopment.

263 **Alternative mechanisms for fate specification and sensory input specificity in projection neurons**

264 What is the origin of signals that govern projection neuron fate and sensory input specificity, if not motor-derived? In comparable
265 systems, fate signals can be intrinsically-expressed or originate from extrinsic sources. For example, intrinsic genetic mechanisms
266 assemble laminar connectivity in visual circuits^{87,88} and facilitate sensorimotor matching in spinal circuits^{9,89-91}. In directionally-
267 selective retinal circuits, subtype fate is established in a similar manner^{92,93}. In “intrinsic” models, synaptic specificity arises from
268 molecular matching between subtypes⁹⁴⁻⁹⁶. Alternatively, in somatosensory and auditory circuits, transcriptional fate depends on
269 extrinsic signals such as growth factors⁹⁷ and sensation⁹⁸, respectively. In spinal circuits, positional fate, which constrains connec-
270 tivity¹³, is established by extrinsic codes such as morphogen gradients in early development⁹⁹ and Hox factors^{63,64}. In “extrinsic”
271 models, early inputs are often erroneous and refined by activity^{100,101} or molecular factors¹⁰². Collectively, these findings offer two
272 alternative models for how vestibulo-ocular reflex circuit assembly emerges.

273 The tight links between birth order, somatic position, and stimulus selectivity^{28,29,33,85,103,104} across vestibulo-ocular reflex circuit
274 populations support an “intrinsic” determination model. Further, neurogenesis and initial axon targeting develops contempora-
275 neously for sensory afferents^{55,103,105,106}, projection neurons^{3,29,107-109}, and extraocular motor neurons^{3,33,39,110-112}, suggest-
276 ing that neurons are poised to assemble with targets as early as their time of differentiation. Importantly, an “intrinsic specification”
277 model makes a testable prediction about how and when sensory selectivity should emerge across the circuit: projection neurons
278 and extraocular motor neurons should be directionally selective as soon as pre-synaptic input is established. Such evidence would
279 justify future molecular inquiries into the underlying genetic factors, expanding early characterizations of the mechanisms that
280 shape hindbrain topography^{113,114}, recent molecular profiling of the zebrafish hindbrain^{115,116}, and reports of molecular match-
281 ing between extraocular motor neurons and muscle^{117,118}. Operationally, the present study lays a foundation for molecular explo-
282 rations of projection neuron subtype determinants by establishing bulk- and single-cell transcriptomic profiling and *in situ* valida-
283 tion pipelines.

284 Conversely, evidence that stimulus selectivity emerges gradually would suggest that sensory afferents and/or projection neu-
285 rons initially wire indiscriminately and that circuit connectivity is refined in time by extrinsic forces. Prior work in the vestibulo-
286 ocular reflex circuit has proposed developmental roles for sensory-derived trophic factors¹¹⁹ and activity-dependent refine-
287 ment^{120,121}, though sensory afferents develop typically in the absence of utricular input¹²² and ocular motor behavior does not

288 depend on stimulus-driven activity¹²³. Here, an “extrinsic” determination model would predict that connectivity is established by
289 an anterogradely-transmitted signal – that is, from sensory afferents to ocular muscles. If so, then future investigations might con-
290 stitutively ablate sensory afferents to eliminate activity-driven, diffusible, or cell-surface instructional signals, similar to the present
291 study. The directional bias in opsin-evoked activity in projection neurons²⁵, together with their transcriptional profiles established
292 here, offer a clear readout of the role of sensory-derived factors. However, genetic targets exclusive to vestibular sensory afferents
293 for gaze stabilization have not been identified, and tissue-specific genetic ablations remain limited in zebrafish. Looking ahead, re-
294 solving when and how stimulus selectivity emerges across the vestibulo-ocular reflex circuit will be key to understanding whether
295 connectivity with pre- and/or post-synaptic partners instructs subtype fate, or whether subtype fate instructs connectivity.

296 **Conclusion**

297 Here, we discovered that motor partners do not determine pre-motor fate and sensory connectivity for the projection neurons that
298 stabilize gaze. Our results overturn the current model that stimulus selectivity and connectivity are retrogradely specified, a major
299 step towards understanding the origin, and eventually nature, of mechanisms that assemble an archetypal sensorimotor reflex cir-
300 cuit. Instead, our data support and extend recent models in spinal systems that motor partners do not actively construct sensory-
301 to-interneuron reflex circuit architecture, but may later refine their inputs. By defining the contribution of motor neurons to specifi-
302 cation and sensory connectivity of gaze-stabilizing central projection neurons, our work speaks to general principles of sensorimotor
303 circuit assembly.

304 **MATERIALS AND METHODS**

305 **RESOURCE AVAILABILITY**

306 **Lead Contact**

307 Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, David
308 Schoppik (schoppik@gmail.com).

309 **Materials Availability**

310 Mutant fish lines generated in this study will be deposited to the Zebrafish International Resource Center (ZIRC).

311 **Data and code availability**

312 • All data and code are deposited at the Open Science Framework and are publicly available at DOI: 10.17605/OSF.IO/93V6E

313 **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

314 **Fish care**

315 All protocols and procedures involving zebrafish were approved by the New York University Langone School of Medicine Institu-
316 tional Animal Care & Use Committee (IACUC). All larvae were raised at 28.5°C at a density of 20-50 larvae in 25-40 ml of buffered
317 E3 (1mM HEPES added). Larvae used for photofill experiments were raised in constant darkness; all other fish were raised on a stan-
318 dard 14/10h light/dark cycle. Larvae for experiments were between 3-5 days post-fertilization (dpf).

319 **Transgenic lines**

320 Experiments were conducted on the *mifta*^{-/-} background to remove pigment. All experiments used larvae from the F3 genera-
321 tion or older of a newly-created line of *phox2a* mutants (described below) on the following backgrounds: *Tg(isl1:GFP)*⁴⁰ to validate
322 *phox2a* loss-of-function; *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16)*^{25,49} to drive UAS reporter expression; *Tg(UAS-E1b:Kaede)*⁴⁸ for
323 anatomical imaging experiments; and *Tg(UAS:GCaMP6s)*⁸³ for calcium imaging experiments. All larvae were selected for bright-
324 ness of fluorescence relative to siblings. Mendelian ratios were observed, supporting that selected larvae were homozygous for fluo-
325 rescent reporter alleles.

326 **Generation of *phox2a* mutants**

327 *phox2a* mutant lines were generated using CRISPR/Cas9 mutagenesis. Two guide RNAs (gRNAs) were designed using the Bench-
328 ling CRISPR Guide RNA Design Tool (see: key resources, [Table 5](#)). gRNAs were located towards the 5' region of exon 1 to minimize
329 the size of any translated protein. gRNAs were incubated with Cas9 protein before co-injection into *Tg(isl1:GFP)* embryos at the
330 single cell stage. Injected embryos were screened for anatomical phenotypes (reduction in *isl1*-positive nIII/nIV motor neurons).
331 Phenotypic embryos (F0) and their embryos were raised and genotyped via sequencing to identify and validate germline muta-
332 tions. Three founders were identified and used for experiments: (1) *phox2a*^{d22} has a 22 bp deletion from base pairs 249 to 270,
333 (2) *phox2a*^{d19} has a 19 bp deletion from base pairs 262 to 280, and (3) *phox2a*ⁱ² has a 2 bp insertion (AG) from base pairs 261 to
334 262. Each mutation created a nonsense mutation, causing a predicted premature stop codon at the beginning of the homeobox.
335 All alleles were validated using complementation assays, and larvae from all three alleles were used in experiments. For brevity, only
336 one allele (*phox2a*^{d22}) is shown in [Figure 1](#).

337 **Maintenance of *phox2a* adults**

338 *phox2a* null larvae do not survive past 7 dpf. Sibling embryos (*phox2a*^{+/+} or *phox2a*^{+/-}) were raised and genotyped to identify het-
339 erozygotes for line propagation. Primers for genotyping are listed in the Key Resources table ([Table 5](#)). Genomic DNA was ampli-
340 fied using a polymerase (DreamTaq PCR Master Mix 2X, Thermo Fisher Scientific K1071), 60° annealing temperature, 30 second
341 elongation time, and 35 cycles of PCR. PCR generates a 169 bp product (wildtype), 147 bp product (*phox2a*^{d22}), 150 bp product
342 (*phox2a*^{d19}), or 171 bp product (*phox2a*ⁱ²). *phox2a*^{d22} and *phox2a*^{d19} DNA was evaluated using gel electrophoresis; *phox2a*ⁱ² was
343 assessed via sequencing with the reverse primer (Genewiz, Azenta Life Sciences, South Plainfield, New Jersey).

344 **METHOD DETAILS**

345 **Confocal imaging**

346 Larvae were anesthetized in 0.2 mg/mL ethyl-3-aminobenzoic acid ethyl ester (MESAB, Sigma-Aldrich E10521, St. Louis, MO) prior
347 to confocal imaging except where noted. Larvae were mounted dorsal side-up (axial view) or lateral side-up (sagittal view) in 2%
348 low-melting point agarose (Thermo Fisher Scientific 16520) in E3. Images were collected on a Zeiss LSM800 confocal microscope
349 with a 20x water-immersion objective (Zeiss W Plan-Apochromat 20x/1.0). Images of tangential nucleus soma and axons were ac-
350 quired in a lateral mount with an 80x80 μm imaging window. Stacks spanned ~30-40 μm, sampled every 1 μm. Images of nIII/nIV
351 motor neurons were acquired in a dorsal mount with a 213x106 μm imaging window; stacks spanned approximately 90 μm, sam-
352 pled every 1.5 μm. Images to validate nIII/nIV expression in a lateral mount were acquired using a 319x319 μm imaging window.
353 Raw image stacks were analyzed using Fiji/ImageJ [124](#).

354 **Identification of *phox2a* larvae**

355 Prior to experiments, larvae were designated as *phox2a* mutants or sibling (wildtype/heterozygote) controls based on two criteria:
356 gross loss of *Tg(isl1:GFP)* fluorescence in nIII/nIV at 2 dpf, visualized using a SugarCube LED Illuminator (Ushio America, Cypress CA)
357 on a stereomicroscope (Leica Microsystems, Wetzlar, Germany) and absence of a swim bladder at 5 dpf. For anatomical and cal-
358 cium imaging experiments, allele designations were validated using confocal imaging of nIII/nIV motor neurons: total or near-total
359 loss of nIII/nIV neurons (null), selective loss of IR/MR neurons (heterozygote), or normal expression (wildtype). Designations were
360 confirmed after experiments using genotyping. For RNA sequencing and fluorescent *in situ* experiments, sibling controls (wild-
361 type/heterozygote) were combined.

362 **Birthdating of nIII/nIV motor neurons**

363 Early-born neurons in nIII/nIV were optically tagged using *in vivo* birthdating [29,33,125](#) on *Tg(isl1:Kaede)^{ch103}* larvae [126](#). Briefly,
364 whole embryos were exposed to UV light for five minutes at experimenter-defined timepoints and subsequently raised in darkness
365 to prevent background conversion. At 5 dpf, larvae were imaged on a confocal microscope. Neurons born before the time of photo-
366 conversion expressed red, converted Kaede; neurons born after expressed only green, unconverted Kaede.

367 **Fluorescent *in situ* hybridization and imaging**

368 Experiments were performed using Hybridization Chain Reaction (HCR) for whole-mount zebrafish larvae^{51,127}. Probes were gen-
369 erated using the HCR 3.0 probe maker¹²⁸ using the sense sequence of the canonical gene cDNA from NCBI. All larvae were from
370 the *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* background. Larvae were pre-identified as null mutants or sib-
371 lings (wildtype or heterozygotes) and combined in equal ratios (8-10 larvae per condition, 16-20 larvae total) into a single 5 mL
372 centrifuge tube for fixation and HCR. Larvae were fixed overnight with 4% PFA in PBS at 4° C and stored in 100% methanol at -
373 20° C. Subsequently, HCR was performed as described in¹²⁷, with adjustments to proteinase K incubation time based on age (3
374 dpf: 30 min incubation; 5 dpf: 50 min incubation). HCR experiments used buffers and amplifiers from Molecular Instruments (Los
375 Angeles, CA). Samples were stored in 1x PBS at 4° C and imaged on a confocal microscope within four days. Prior to imaging, larvae
376 were re-screened for *Tg(isl1:GFP)* fluorescence to identify null mutants and sibling controls. For each probe, imaging parameters
377 were determined using a sibling control and kept constant for all subsequent larvae. Comparable settings (within 1% laser power)
378 were used across probes.

379 **Calcium imaging of tonic and impulse tilt stimuli responses**

380 Experiments were performed as described in²⁹ using Tilt-In-Place Microscopy⁴¹. All experiments used 5 dpf larvae from the
381 *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:GCaMP6s)* background. Briefly, larvae were mounted dorsal-up in 2% low-melt
382 agarose in E3 onto a large beam diameter galvanometer system (ThorLabs GVS011). Tonic pitch-tilt stimuli were presented over a
383 65-second period in the following order: horizontal baseline (5 sec at 0°), nose-down tilt (15 sec at -19°), horizontal imaging (15 sec
384 at 0°), nose-up tilt (15 sec at 19°), and horizontal imaging (15 sec at 0°). Impulse stimuli contained a 4 msec eccentric rotation, a
385 2 msec hold, and a 4 msec restoration step to horizontal and were presented twice over a 65-second imaging window: horizontal
386 baseline (20 sec), impulse (10 msec), horizontal imaging (30 sec), impulse (10 msec), horizontal imaging (15 sec). Tonic and im-
387 pulse stimuli were presented in alternating sets (impulse, then tonic) with a total of three stimulus set repeats.

388 Imaging was performed using a 20x water immersion objective (Olympus XLUMPLFLN20xW 20x/1.0), an infrared laser (Spectra-
389 Physics MaiTai HP) at 920nm using 6.1-18.8 mW of power at the sample, and ThorLabs LS 3.0 software. Experiments were con-
390 ducted in the dark. High-resolution anatomy scans of nIII/nIV motor neurons were performed for each experiment to validate allele
391 designations. Scans used a 147x147 µm imaging window, a 90 µm stack sampled every 1.5 µm, and a 5.2 microsecond pixel dwell
392 time. Anatomy scans of the tangential nucleus were acquired using a 148x91 µm imaging window as a 40-50 µm stack sampled
393 every 1 µm. For stimulus imaging, the tangential nucleus was sampled every 3-6 µm based on cell density. 6-10 planes were sam-
394 pled for each hemisphere. Ventral planes were imaged at higher magnification (112x68 µm imaging window) than dorsal planes
395 (148x91 µm window) to avoid photomultiplier tube saturation from in-frame GFP fluorescence; magnification was corrected for in
396 later analyses. Laser power was adjusted for each sampled plane due to the light scattering properties of zebrafish tissue. As greater
397 power was required for ventral planes, imaging was always performed from ventral to dorsal to minimize photobleaching effects.
398 Stimulus imaging was performed at 3 frames/second (2.2 µs pixel dwell time) with a total time of approximately two hours per fish.

399 **Retrograde photolabeling of tangential nucleus neurons**

400 Experiments were performed as described in²⁹ based on⁴⁴ on 5 dpf larvae from the *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-*
401 *VP16);Tg(UAS-E1b:Kaede)* background. Briefly, experiments leveraged a photoconvertible protein, Kaede, which irreversibly con-
402 verts from green to red with ultraviolet light. Larvae were raised in darkness to minimize background conversions. Larvae were
403 mounted dorsal-up in 2% agarose under a confocal microscope. An imaging window was centered over the medial longitudinal
404 fasciculus (MLF) and repeatedly scanned with a 405 nm laser for 30 seconds until fully converted (green to red). Off-target photo-
405 conversion was assessed (e.g., conversion of projections lateral to the MLF). Larvae were unmounted, left to recover in E3 for 4 hours
406 in darkness, and then re-mounted in a lateral mount. An imaging window was centered around the tangential nucleus (see: Con-
407 focal Imaging). Retrogradely-labeled soma were identified by their center-surround fluorescence appearance: red converted cyto-
408 plasm surrounding an unconverted green nucleus.

409 **Neuron harvesting, dissociation, and flow cytometry**

410 Experiments were performed on 72-74 hpf larvae from the *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* back-
411 ground. At 2 dpf, larvae were designated as null or sibling (wildtype/heterozygote) as described above. Three experimenters (D.G.,
412 K.R.H., and P.L) harvested neurons in parallel. Larvae were anesthetized in MESAB in Earle's Balanced Salt Solution with calcium,
413 magnesium, and phenol red (EBSS, Thermo Fisher Scientific 24010043) and mounted dorsal-up in 2% agarose. Fluorescence in
414 tangential nucleus neurons was visualized using a SugarCube LED Illuminator (Ushio America, Cypress CA) using 10x eyepieces on
415 a stereomicroscope (Leica Microsystems, Wetzlar, Germany). Neurons were harvested using a thin wall glass capillary tube (4 inch,
416 OD 1.0 MM, World Precision Instruments) into EBSS in a non-stick Eppendorf tube and kept on ice until dissociation.

417 Neurons were dissociated in 20 units/mL of papain prepared in EBSS (Worthington Biochemical), 2000 units/mL of deoxyribonu-
418 cleic prepared in EBSS (Worthington Biochemical), and 100 mg/mL of Type 1A Collagenase (Sigma Aldrich) prepared in Hanks
419 Buffered Salt Solution without calcium/magnesium (HBSS, Thermo Fisher Scientific). Neurons were incubated for 45 minutes at
420 31.5°C with a gentle vortex every 10-15 min, then passed through a 20 µm filter and centrifuged for 10 mins at 300 x g. After re-
421 moving supernatant, neurons were resuspended in L15 (Thermo Fisher Scientific) with 2% fetal bovine serum (Thermo Fisher Sci-
422 entific). Cell health was evaluated using DAPI, applied at 0.5 µg/ml (Invitrogen) and incubated on ice for 30-45 mins prior to flow
423 cytometry.

424 Flow cytometry was performed using a Sony SH800z cell sorter (100 µm nozzle, 20 psi) to isolate single neurons (Figure S2). Three
425 controls were run: (1) non-fluorescent wildtype neurons, (2) non-fluorescent neurons + DAPI, (3) fluorescent (green) neurons from
426 *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* + DAPI. On average, 2% of neurons were DAPI-positive and ex-
427 cluded. Neurons were evaluated for positive (green) fluorescence. Fluorescence was not evaluated to separate *Tg(UAS-E1b:Kaede)*
428 neurons from those labeled by *Tg(isl1:GFP)*. Neurons were sorted into an Eppendorf tube containing 700 µl of lysis buffer (RNAque-
429 ous Micro Total RNA Isolation Kit, Thermo Fisher Scientific) for downstream bulk RNA sequencing.

430 **Bulk RNA sequencing**

431 RNA isolation was performed using an RNAqueous Micro Total RNA Isolation Kit (Thermo Fisher Scientific). RNA concentration and
432 quality (RIN > 8.0) was evaluated using an RNA 6000 Pico Kit and a 2100 Bioanalyzer system (Agilent Technologies, Santa Clara,
433 California). RNA sequencing was performed by the NYU Genome Technology Center. Libraries were prepared using the low-input
434 Clontech SMART-Seq HT with Nxt HT kit (Takara Bio USA) and sequenced using an Illumina NovaSeq 6000 with an S1 100 Cycle
435 Flow Cell (v1.5).

436 **QUANTIFICATION AND STATISTICAL ANALYSIS**

437 **Cell counting and spatial mapping of nIII/nIV motor neurons**

438 Analysis was performed in Fiji/ImageJ¹²⁴ using the Cell Counter plugin. Anatomical stacks of nIII/nIV were subdivided in the
439 dorsoventral axis as described in³³ to facilitate localization. A point ROI was dropped over each neuron in the plane in which the
440 soma was brightest (center). The number of neurons in each dorsoventral plane and their coordinates were recorded. Neuron co-
441 ordinates were standardized relative to a (0,0) point, defined as one corner of a standard-sized rectangular box centered over the
442 extent of nIII/nIV in a maximum intensity projection. Differences in spatial location across genotypes was evaluated separately for
443 each spatial axis using a two-tailed, two-sample Kolmogorov-Smirnov test. Probability distributions for figures were generated using
444 the mean and standard deviation from bootstrapped data (n=100 iterations) to ensure results were robust to data from single larva.

445 **Analysis of calcium imaging experiments**

446 Analysis methods are detailed in²⁹ and summarized briefly here. Regions of Interest (ROIs) were drawn around tangential nucleus
447 neurons for each stimulus plane sampled and adjusted for minor movement (1-2 µm) between trials. Raw fluorescence traces were
448 extracted using Matlab R2020b (MathWorks, Natick, Massachusetts) and normalized by ROI size to account for variation in magni-
449 fication. A neuron's response to tonic or impulse stimuli was defined as the change in fluorescence in the first second of restoration

450 to horizontal following tilt delivery. Responses were normalized using a baseline period, defined as the mean fluorescence across
451 the initial baseline window (5 sec) preceding the nose-down tilt (nose-down response) or the last 3 sec of the horizontal restoration
452 following nose-down tilt (nose-up response). This was used to generate a ΔFF value. A ΔFF response was defined as significant if it
453 was greater than two standard deviations above baseline. Directional selectivity was assigned by normalizing the difference in ΔFF
454 responses to each tilt by their sum. This generated a scale of values of ± 1 (i.e., positive values represent nose-up selectivity; negative
455 values, nose-down). Some neurons responded to both tilt directions with high similarity; we set a minimum threshold of $\text{abs}(0.1)$ to
456 distinguish neurons with a clear directional selectivity from untuned neurons.

457 **Spatial mapping of tangential nucleus neurons**

458 Analysis methods are detailed in²⁹ and summarized briefly here. All imaged neurons were manually registered to a reference
459 framework using Adobe Illustrator (2021). Anatomy stacks from all experiments were aligned in the XY (rostr-caudal, mediolat-
460 eral) axes using established anatomical landmarks (e.g., Mauthner cell body, medial longitudinal fasciculus, otic capsule). For Z-
461 registration (dorsoventral axis), stacks were subdivided into eight sections using landmarks within and around the tangential nu-
462 cleus (e.g., Mauthner cell body, neuropil). All registered images were verified by two independent observers (D.G. and S.H.). Neurons
463 were localized to one dorsoventral section and a reference circle, representing a cell, was placed in Illustrator. Coordinates for each
464 reference circle were recorded and standardized to an absolute (0,0) point (dorsomedial-most point of the tangential nucleus). Co-
465 ordinates were imported into Matlab (R2020b) and used to generate a spatial map of imaged neurons.

466 **Statistical analysis of differences in tilt responses across *phox2a* genotypes**

467 Statistical comparisons of tonic and impulse tilt responses are summarized in Table 1. Analyses used a one-way analysis of variance
468 with multiple comparisons. No significant differences (tonic tilt responses) or small differences (impulse responses) were observed
469 across genotypes. Control data reported in Results and Figure 2-Figure 3 is an aggregate from wildtype, *phox2a*^{+/+}, and *phox2a*^{+/-}
470 larvae.

471 **Alignment, quality control, and differential expression analysis of bulk sequencing data**

472 Initial alignment and analyses were performed by the Applied Bioinformatics Laboratories at the NYU School of Medicine
473 (RRID:SCR_019178). Sequencing data was aligned to the GRCz11 zebrafish reference genome and two fluorescent markers
474 (Kaede, GFP; NCBI). Eight datasets from four experimental repeats were aligned: four from *phox2a* mutants, and four from sib-
475 ling controls. One experimental repeat had significantly higher variance in the first and second principal components, likely due to
476 poor quality leading to extremely low transcript counts, and was excluded from downstream analyses. Number of cells/larvae se-
477 quenced and used in downstream analysis are as follows: Repeat 1, n=532/n=904 cells from N=28/N=28 *phox2a* null/control lar-
478 vae; Repeat 2, n=802/n=683 cells from N=27/N=26 *phox2a* null/control larvae; Repeat 3, n=1000/n=1007 cells from N=41/N=40
479 *phox2a* null/control larvae; Repeat 4 (excluded): n=690/n=571 cells from N=33/N=33 *phox2a* null/control larvae. Differential gene
480 expression between conditions (*phox2a* mutants vs. sibling controls) was assessed using DESeq2¹²⁹. Differentially-expressed can-
481 didate genes met two criteria: \log_2 fold change $>|2|$ and p adjusted < 0.05 .

482 **Filtering of bulk sequencing data using a reference single-cell sequencing dataset**

483 Analyses were performed in R. Detection of markers for motor neurons (*isl1*, *isl2a*, *isl2b*)^{130,131} and neurons caudal (*hoxd4a*)^{113,132}
484 and lateral (*barhl2*)¹¹⁴ to rhombomeres 4-6 supported that our dataset included other populations. We applied a filter to exclude
485 erroneous gene expression from non-tangential nucleus populations.

486 Filtering was performed using an existing single-cell atlas of neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);(Tg(UAS-E1b:Kaede)*,
487 generated with 10x Genomics. The reference atlas was generated from four experimental samples using the harvest, dissociation,
488 and flow cytometry method described above. The sequenced atlas contained 1,468 neurons (Figure S3A-Figure S3B)). Data was
489 analyzed using Seurat v4.0¹³³. Cluster annotation was performed using a combination of fluorescent *in situ* hybridization as de-
490 scribed above (Figure S3C-Figure S3E and other data not shown) and published molecular data of the zebrafish hindbrain¹¹³.

491 n=473 neurons (32%) were validated as excitatory projection neurons from the tangential nucleus.

492 Genes in the bulk dataset were only included in downstream analyses if they were expressed above threshold percent of reference
493 projection neurons: 1%, 3%, 5%, 10%, 30%, or 50%. The most stringent filter (50%) was set using the transcription factor *evx2*, which
494 is reported to be expressed in all tangential nucleus neurons⁵⁰ and was detected in 50% of reference projection neurons. Qualita-
495 tively, we found that gene detection with fluorescent *in situ* hybridization scaled with reference filter stringency (Figure S4). Anal-
496 yses were performed separately for each threshold. The total number of genes included for downstream analyses for each thresh-
497 old are as follows: 28,807 (no threshold), 11,189 (1% of reference neurons), 7,871 (3%), 6,075 (5%), 3,579 (10%), 818 (30%), 288
498 (50%). We used the following significance thresholds for differential gene expression in filtered datasets: adjusted p value < 0.05
499 and $\text{abs}(\log_2\text{FoldChange}) > 2$. The number of differentially expressed genes for each threshold was as follows: 91 (no threshold),
500 14 (1% of reference neurons), 3 (3%), 2 (5%), 0 (10%).

501 Projection neurons in the tangential nucleus are transcriptionally similar to excitatory neurons in the medial vestibular nucleus
502 (MVN; unpublished data). MVN neurons may be included in our bulk sequencing dataset given their exceptionally close proximity
503 (3-5 μm) to the medial edge of the tangential nucleus. Some MVN neurons express *phox2a* (Figure S6). To control for the possibility
504 that some differentially expressed genes are localized to the MVN, and not projection neurons, we also evaluated differential gene
505 expression in a validated subset of excitatory MVN neurons (n=271 neurons; 18% of reference dataset) from the same single-cell at-
506 las. Data is shown in Figure S6.

507 Generation of representative images for fluorescent *in situ* hybridization

508 Images were generated using Fiji/ImageJ¹²⁴. An anatomical template of the tangential nucleus was generated based on²⁹. Briefly,
509 for sagittal view images, a 30- μm stack was centered over the tangential nucleus. For each plane, a region of interest (ROI) was
510 drawn over all cells within the bounds of the tangential nucleus. Transcript expression outside the ROI was masked. Maximum in-
511 tensity projections were generated. Minimal or no alterations to brightness/contrast were made for probe expression given the cor-
512 relation between fluorescence intensity and detected transcript⁵¹.

513 Additional statistics

514 Bias and variability in probability distributions were estimated by bootstrapping, or resampling the raw distributions with replace-
515 ment¹³⁴. Data shown is the mean and standard deviation of 100 bootstrapped distributions. Topography data was evaluated using
516 two-tailed, two-way Kolmogorov-Smirnov tests. Functional responses to tilts (i.e., calcium response strength, directionality index)
517 were evaluated using two-tailed Wilcoxon rank sum tests. Differences in responses across genotypes were analyzed using one-way
518 analysis of variance tests.

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AUTHOR CONTRIBUTIONS

Conceptualization: DG and DS, Methodology: DG, BR, KRH, PL, and DS, Investigation: DG, BR, KRH, PL, ML, HP, HG, SH, and CQ, Visualization: DG, Writing: DG, Editing: DS, Funding Acquisition: DG and DS, Supervision: DS.

AUTHOR COMPETING INTERESTS

The authors declare no competing interests.

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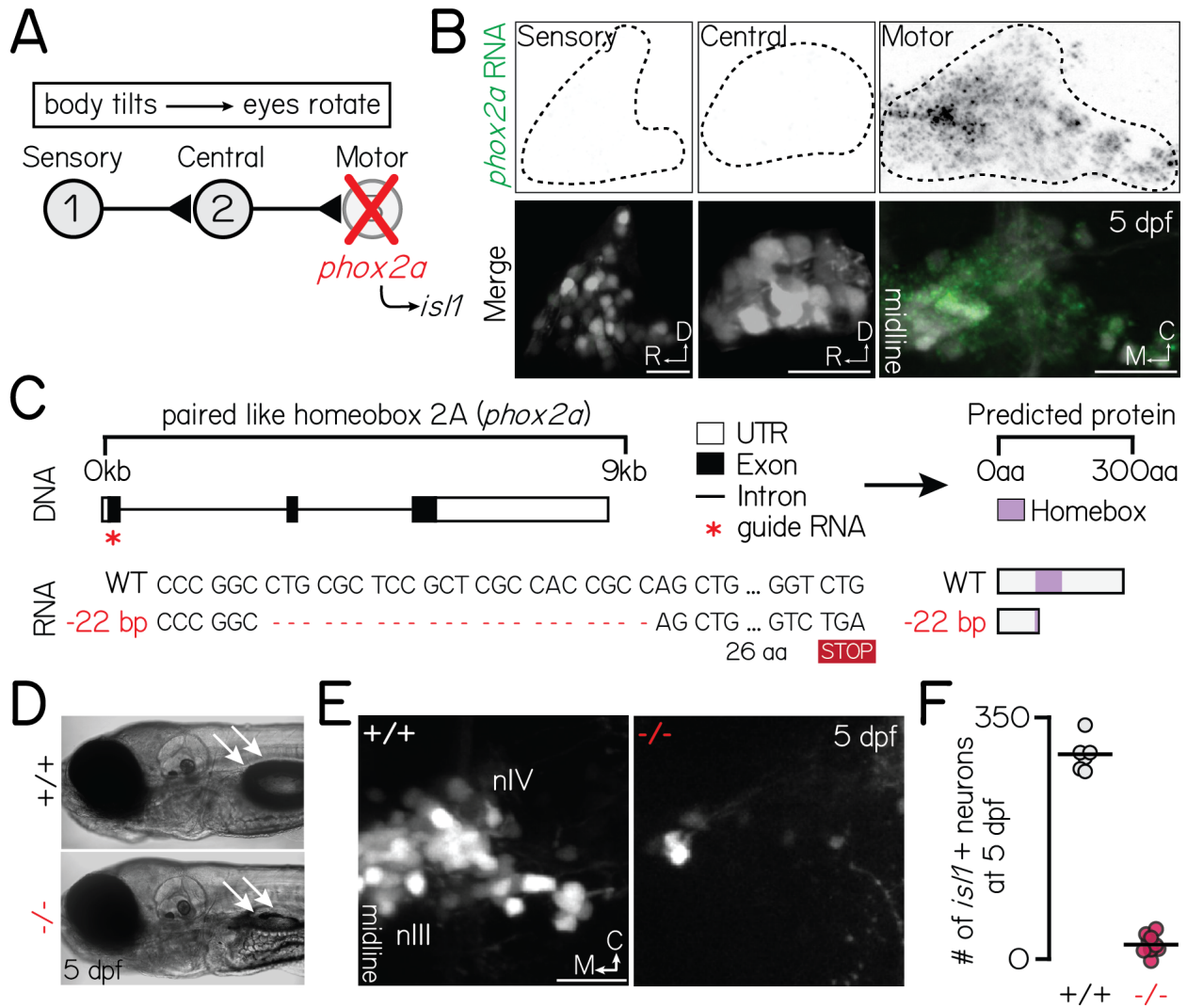


Figure 1: *phox2a* loss-of-function mutants fail to develop nIII/nIV motor neurons and vertical eye rotation behavior.

Associated with [Figure S1](#).

(A) Schematic of vestibulo-ocular reflex circuitry and the genetic loss-of-function approach used to perturb motor-derived signals.

(B) Fluorescent *in situ* hybridization showing *phox2a* transcript expression in statoacoustic ganglion sensory afferents (left), central projection neurons in the tangential nucleus (middle), or nIII/nIV extraocular motor neurons (right) at 5 days post-fertilization (dpf). Top: probe only, nuclei outlined with dashed lines. Bottom: probe (green) merged with somata, labeled by *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (sensory, central) or *Tg(isl1:GFP)* (motor).

(C) Schematic of CRISPR/Cas9 mutagenesis approach. Top: Red star shows location of guides against *phox2a* DNA. Bottom: RNA sequence in wildtype and *phox2a*^{d22} alleles. Red dashed lines show deleted sequence; "STOP" box shows predicted premature stop codon due to deletion. Right shows predicted protein sequence.

(D) Transmitted light image of a 5 dpf wildtype (top) and *phox2a* null mutant (bottom). White arrows point to a normally inflated (top) or absent (bottom) swim bladder.

(E) Images of nIII/nIV motor neurons in one hemisphere, labeled by *Tg(isl1:GFP)*, in wildtype siblings (left) and *phox2a* null mutants (right) at 5 dpf. Scale bar, 20 μ m.

(F) Quantification of the number of *Tg(isl1:GFP)*+ neurons in nIII/nIV from N=6 wildtype siblings and N=10 *phox2a* null mutants.

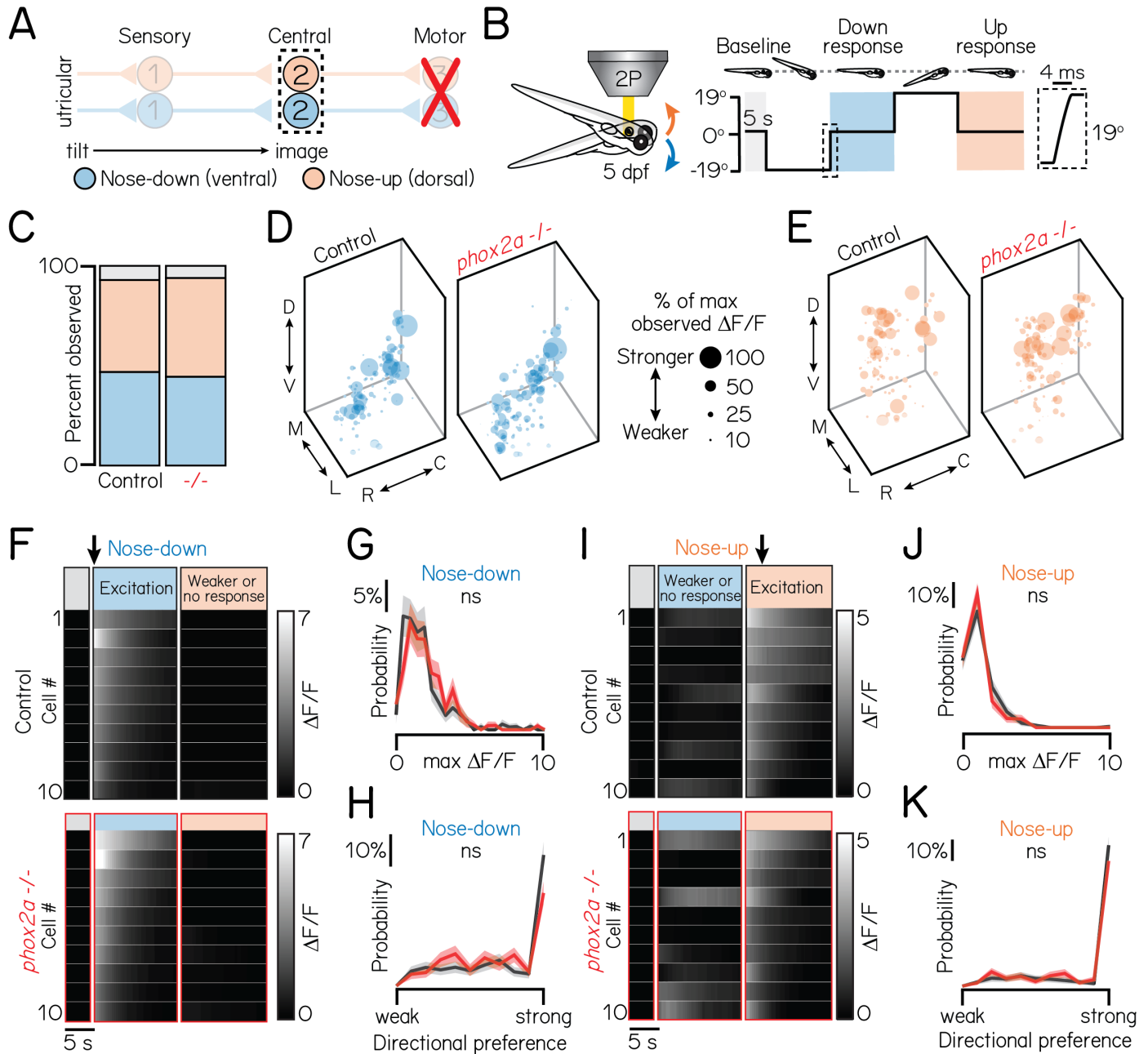


Figure 2: Motor neurons are dispensable for proper connectivity between utricular sensory afferents and projection neurons.

Associated with Table 1.

(A) Schematic of pitch vestibulo-ocular reflex circuitry. Dashed lines outline projection neurons as calcium imaging target. Nose-down/eyes-up channel represented with blue; orange, nose-up/eyes-down.

(B) Schematic of tonic pitch-tilt stimulus delivered with Tilt-In-Place Microscopy (TIPM). Shaded regions show calcium imaging windows when fish were oriented horizontally immediately following tilts. Inset shows timecourse of the rapid step to restore horizontal position after tilts. Imaging experiments used larvae from the *Tg(isl1:GFP);Tg(-6.7Tru.Hcrt2:GAL4-VP16);Tg(UAS:GCaMP6s)* line.

(C) Proportion of subtypes observed in sibling controls and *phox2a* null mutants. Blue: nose-down. Orange: nose-up. Grey: Neurons without directional tuning (criteria in Methods).

(D/E) Soma position of nose-down (blue) and nose-up (orange) neurons in sibling controls (left) and *phox2a* null mutants (right). Soma size scaled by strength of calcium response ($\Delta F/F$), normalized by max observed $\Delta F/F$.

(F/I) Heatmaps showing example tilt responses from nose-down (F) or nose-up (I) neurons in sibling controls (top) and *phox2a* null mutants (bottom). $n=10$ neurons with strongest $\Delta F/F$ responses to tilts shown. Each row shows an individual neuron. Shaded bars show calcium imaging window immediately following restoration from eccentric position. Black arrow points to first second of tilt response used for analyses.

(G/J) Distributions of maximum $\Delta F/F$ responses to tilts for nose-down (G) or nose-up (J) neurons in sibling controls (black) and *phox2a* null mutants (red). Solid and shaded lines show mean and standard deviation, respectively, of bootstrapped data (Methods)

(H/K) Distributions of directional tuning score to tilts for nose-down (H) or nose-up (K) neurons in sibling controls (black) and *phox2a* null mutants (red). Tuning score ranges from 0 (equal responses to both tilt directions, no tuning) to 1 (responses to one tilt direction only); criteria detailed in Methods. Solid and shaded lines show mean and standard deviation, respectively, of bootstrapped data.

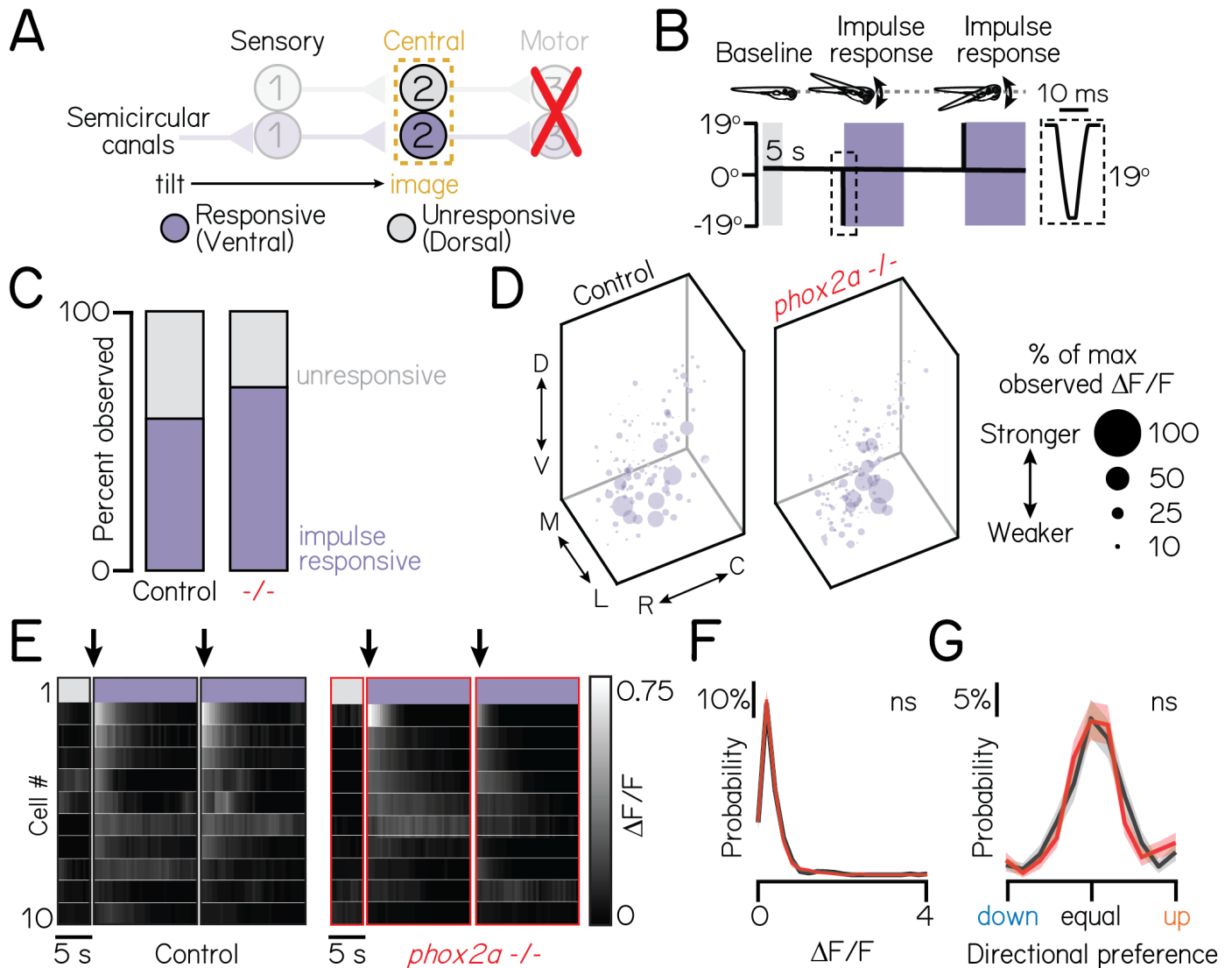


Figure 3: Motor neurons are dispensable for proper connectivity between semicircular canal sensory afferents and projection neurons.

Associated with Table 1.

- (A) Schematic of impulse tilt experiments. Yellow dashed lines outline projection neurons as calcium imaging target. Impulse-responsive neurons (ventrally-localized) shown with purple; unresponsive neurons, grey.
- (B) Schematic of impulse stimuli delivered with TIPM. Shaded regions show calcium imaging windows at horizontal immediately following impulses. Inset shows timecourse of impulse stimulus. Imaging experiments used larvae from the *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:CCaMP6s)* line.
- (C) Proportion of impulse-responsive (purple) and unresponsive (grey) neurons observed in sibling controls and *phox2a* null mutants.
- (D) Soma position of impulse-responsive neurons in sibling controls (left) and *phox2a* null mutants (right). Soma size scaled by strength of calcium response (ΔFF), normalized by max observed ΔFF .
- (E) Heatmaps showing example impulse responses from neurons in sibling controls (left) and *phox2a* null mutants (right). $n=10$ example neurons shown. Each row shows an individual neuron. Shaded bars show calcium imaging window immediately following impulse delivery. Black arrow points to first second of tilt response used for analyses. Note smaller scale (0-0.75) of impulse responses relative to Figures 2F and 2I.
- (F) Distributions of maximum ΔFF responses to impulses in sibling controls (black) and *phox2a* null mutants (red). Solid and shaded lines show mean and standard deviation, respectively, from bootstrapped data.
- (G) Distributions of directional tuning score to impulses in sibling controls (black) and *phox2a* null mutants (red). Tuning score ranges from 0 (equal responses to both tilt directions, no tuning) to 1 (responses to one tilt direction only); criteria detailed in Methods. Solid and shaded lines show mean and standard deviation, respectively, from bootstrapped data.

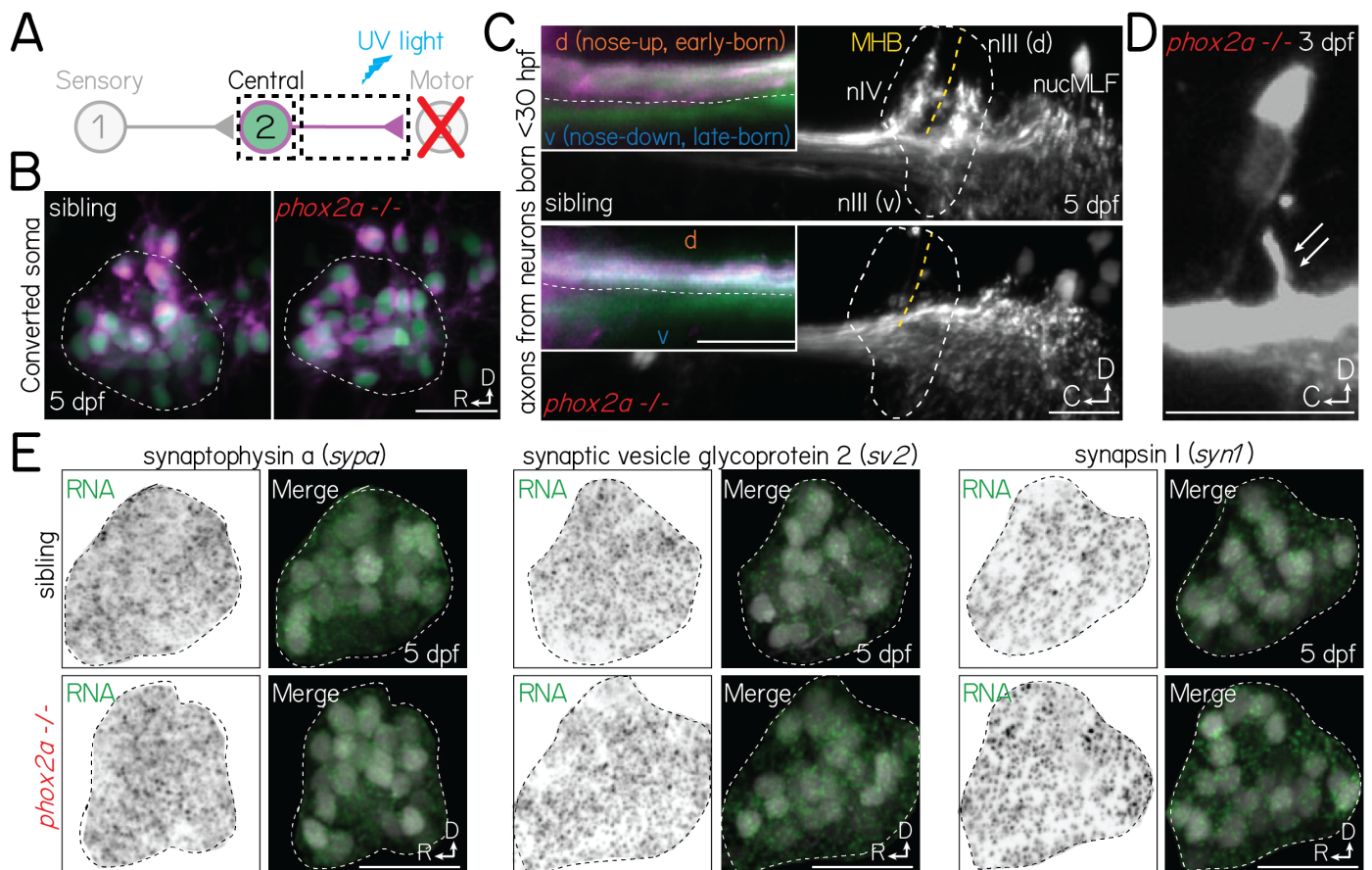


Figure 4: Projection neurons are anatomically and molecularly poised to assemble with motor neuron partners in *phox2a* mutants.

(A) Schematic of retrograde photofill experiments. Projection neuron axons expressing the photolabile protein Kaede are targeted at the midbrain-hindbrain boundary with ultraviolet light. Converted protein (magenta) retrogradely diffuses to the soma, while the unconverted nucleus remains green.

(B) Projection neuron somata in sibling controls (left) and *phox2a* null mutants (right) after retrograde photolabeling. Experiments performed at 5 dpf. Neurons visualized in *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:E1b-Kaede)*.

(C) Projection neuron axons at the hindbrain (inset) and midbrain-hindbrain boundary in sibling controls (top) and *phox2a* null mutants (bottom). Axons visualized using *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:E1b-Kaede)*. White dashed outline shows arborization fields in nIII/nIV. MHB and yellow dashed line, midbrain-hindbrain boundary. nucMLF: nucleus of the longitudinal fasciculus. Inset: Spatial segregation between early-born (magenta+green) and late-born (green only) axons. White dashed line reflects separation between dorsal (nose-up, early-born) and ventral (nose-down, late-born) axon bundles. Image at 5 dpf in sagittal view.

(D) Projection neuron axon bundle in a *phox2a* null mutant at 3 dpf. White arrows point to single collateral to two remaining nIII/nIV neurons.

(E) Fluorescent *in situ* hybridization against RNA for three pre-synaptic markers: synaptophysin a (*sypa*; left), synaptic vesicle glycoprotein 2 (*sv2*, middle), and synapsin I (*syn1*, right). Top row, sibling controls. Bottom row, *phox2a* null mutants. For each panel set, left images show *in situ* probe expression (green) and right images show merge with projection neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:E1b-Kaede)*. Dashed lines outline the projection nucleus. Cell and transcript expression outside the projection nucleus is removed for visual clarity. Images taken at 5 dpf in sagittal mount. All scale bars, 20 μ m.

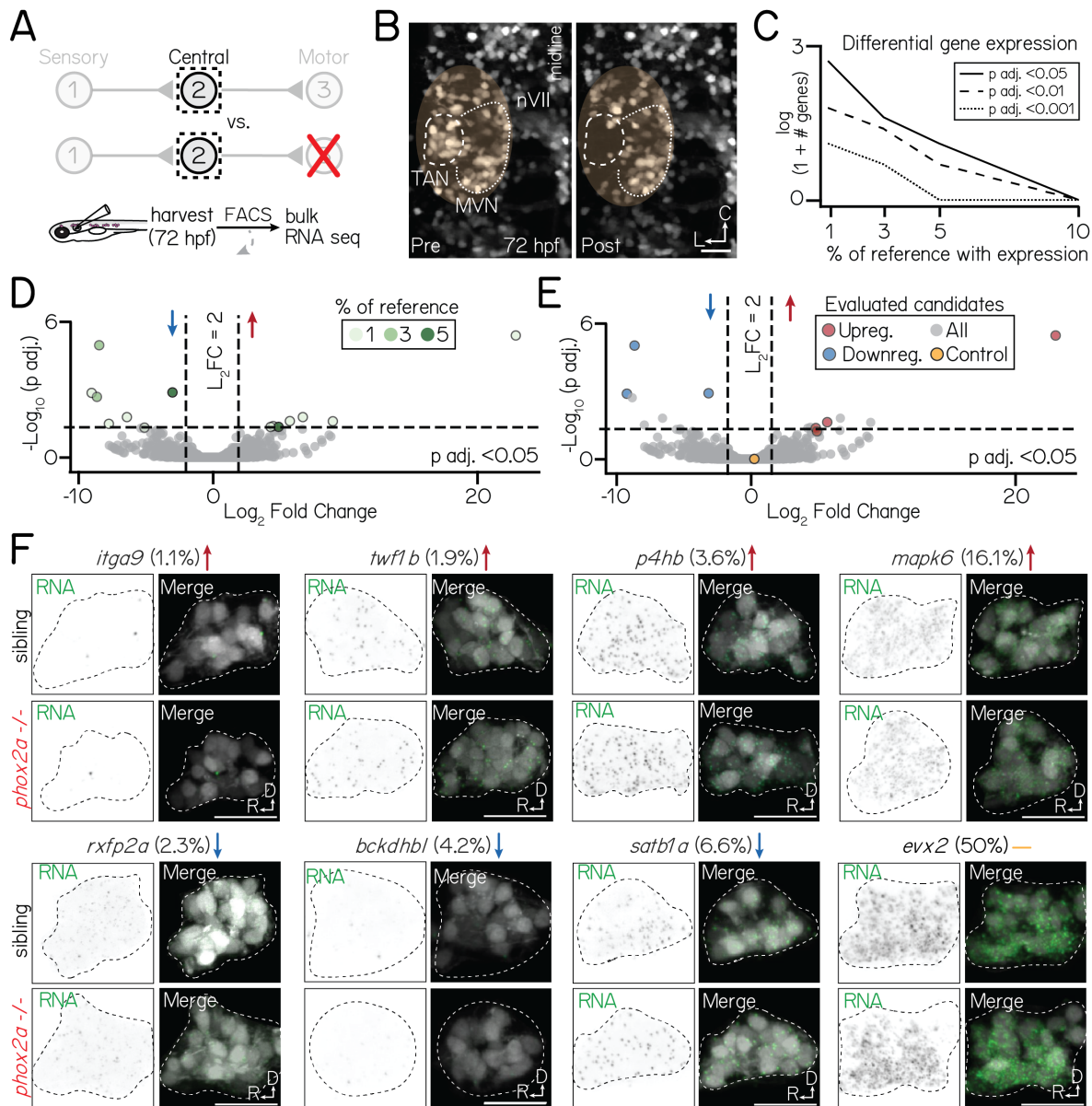


Figure 5: Motor neurons are dispensable for normal transcriptional profiles of projection neurons.

Associated with [Figure S2-Figure S5, Table 2.](#)

(A) Schematic of sequencing approach. Central projection neurons (*Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:E1b-Kaede)*) are harvested from 3 dpf larvae. Flow cytometry is used to exclude neurons not labelled by *Tg(-6.7Tru.Hcrtr2:GAL4-VP16)*. Bulk RNA sequencing is performed to compare the profiles of projection neurons in siblings and *phox2a* null mutants.

(B) Example of projection neurons before (left) and after (right) harvesting. Neurons visualized with *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS:E1b-Kaede)*. Dashed lines outline projection neurons in the tangential nucleus; dotted lines, medial vestibular nucleus. Yellow region shows margin of harvesting error: non-projection neurons that may be included in bulk sequencing dataset.

(C) Number of differentially expressed genes in projection neurons at 3 dpf after applying progressive filters based on gene expression in a reference single-cell dataset. Data shown on logarithmic scale. Solid, dashed, and dotted lines represent differentially-expressed gene with p adjusted < 0.5, p adjusted < 0.01, or p adjusted < 0.001 significance, respectively.

(D) Volcano plot showing differentially expressed genes in projection neurons between control and *phox2a* null larvae at 3 dpf. Dashed lines represent significance cutoffs: horizontal line, p adjusted > 0.05; vertical line, Log_2 Fold Change > 2.0. Each circle is a gene. Genes to the left and right of 0 on the horizontal axis show downregulated and upregulated genes, respectively. Colors indicate percent of reference cells that express a given gene. Grey-colored genes are below both significance thresholds.

(E) Same data as [Figure 5D](#). Colored genes show eight candidates evaluated with fluorescent *in situ* hybridization: red, upregulated; blue, downregulated; yellow, highly-expressed controls (*evx2*).

(F) Fluorescent *in situ* hybridization against candidate genes that met projection neuron filter criteria. Top row shows sibling controls; bottom row, *phox2a* null mutants. For each gene, left panels show RNA probe (green) and right panels show merge with projection neurons labeled by *Tg(-6.7Tru.Hcrtr2:GAL4-VP16)* (grey). Dashed lines outline the projection nucleus. Cell and transcript expression outside the projection nucleus is masked for visual clarity. Arrows denote whether genes are upregulated (red), downregulated (blue), or not significantly changed (yellow). Percentage refers to fraction of cells in a single-cell RNA sequencing reference atlas (Methods) with detected transcript. Candidates: *itga9* (\log_2 fold change = 23.0, p adj. = 3.9×10^{-6}), *twf1b* (\log_2 fold change = 5.9, p adj. = 0.024), *p4hb* (\log_2 fold change = 5.1, p adj. = 0.04), *mapk6* (\log_2 fold change = 5.1, p adj. = 0.06), *rxfp2a* (\log_2 fold change = -8.5, p adj. = 1.1×10^{-5}), *bckdhbl* (\log_2 fold change = -9.1.0, p adj. = 0.001) *satb1a* (\log_2 fold change = -3.0, p adj. = 0.001), *evx2* (\log_2 fold change = 0.46, p adj. = 0.99). All scale bars, 20 μ m.

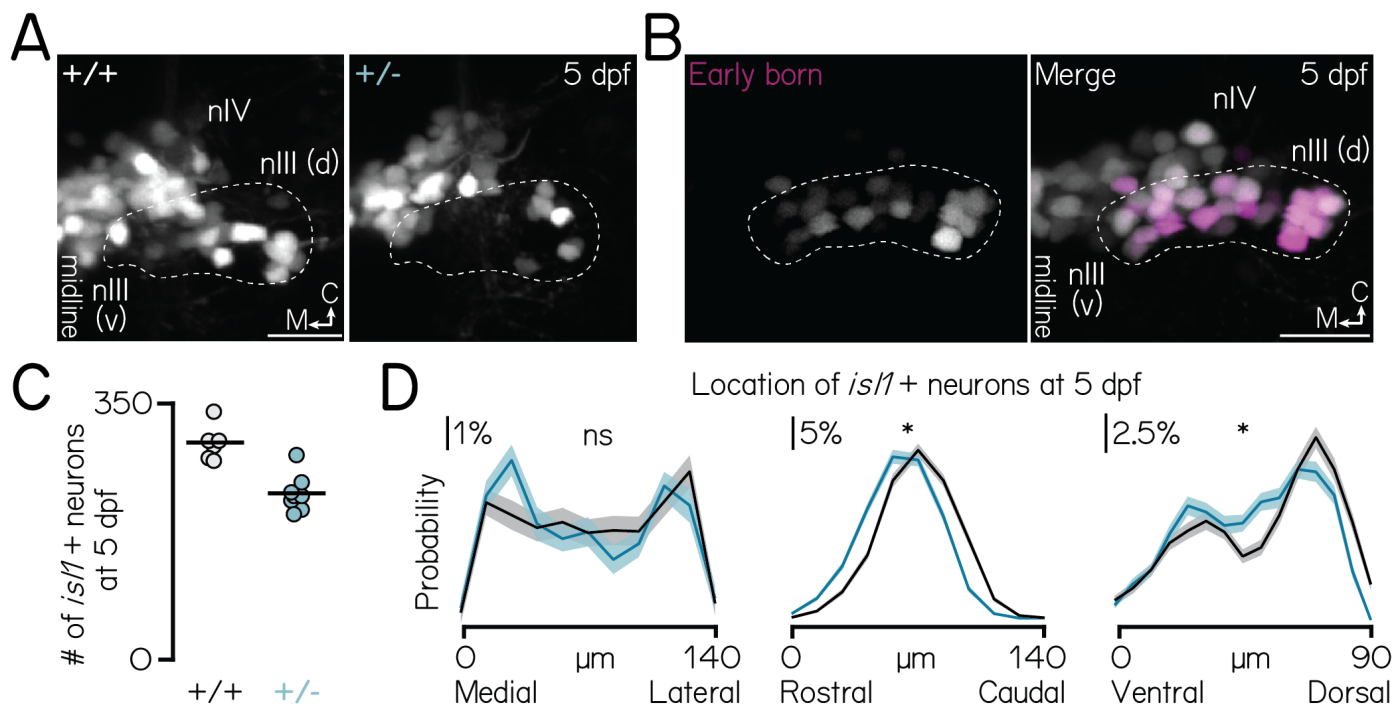


Figure S1: *phox2a* specifies nIII motor neuron fate in a dose- and birthdate-dependent manner.

Associated with Figure 1.

(A) Images of nIII/nIV motor neurons, labeled in *Tg(is1:GFP)*, in wildtype siblings (left) and *phox2a* heterozygotes (middle) at 5 dpf. Wildtype image same as in Figure 1E. One hemisphere shown. White dashed lines outline the dorsal extent of nIII, which contains inferior rectus and medial rectus neurons³³. Scale bar, 20 μ m.

(B) Location of the earliest-born neurons in nIII/nIV (left, magenta) against all nIII/nIV neurons labeled in *Tg(is1:Kaede)* (right, grey). Larvae birthdated at 34 hpf (Methods). One hemisphere shown. White dashed lines outline the dorsal extent of nIII. Scale bar, 20 μ m.

(C) Quantification of the number of *Tg(is1:GFP)*+ neurons in nIII/nIV from N=6 wildtype siblings (grey) and N=8 *phox2a* heterozygotes (teal). Wildtype data same as Figure 1F.

(D) Distributions showing probability of nIII/nIV soma location across each spatial axis in wildtype (black) and heterozygous (teal) *phox2a* larvae. Solid and shaded lines show mean and standard deviation, respectively, from bootstrapped data. Data from same fish quantified in Figure S1C. ns, not significant; star, significant at the $p < 0.001$ level.

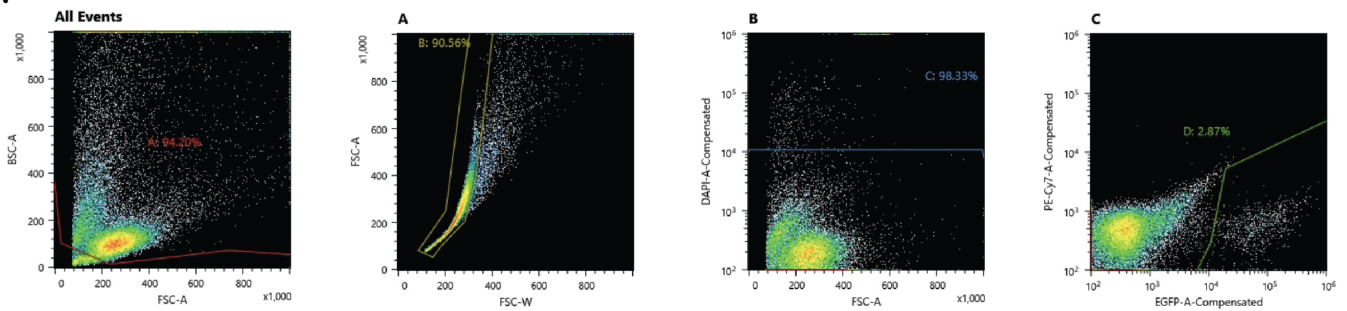
A

Figure S2: Flow cytometry gating strategy to sort fluorescently-labeled neurons for bulk RNA sequencing.

Associated with [Figure 5](#).

(A) Sequential gates used to sort fluorescent neurons labeled with *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede);Tg(isl1:GFP)*. Gate A excluded presumptive debris (small cells). Gate B isolated single cells and excluded large cells and doublets. Gate C excluded DAPI+ (dead or unhealthy) neurons. Gate D isolated fluorescent (GFP or Kaede+) neurons; neurons in this gate were sorted. Gates were set using negative controls (not shown; Methods). Gates shown for one of four experimental repeats.

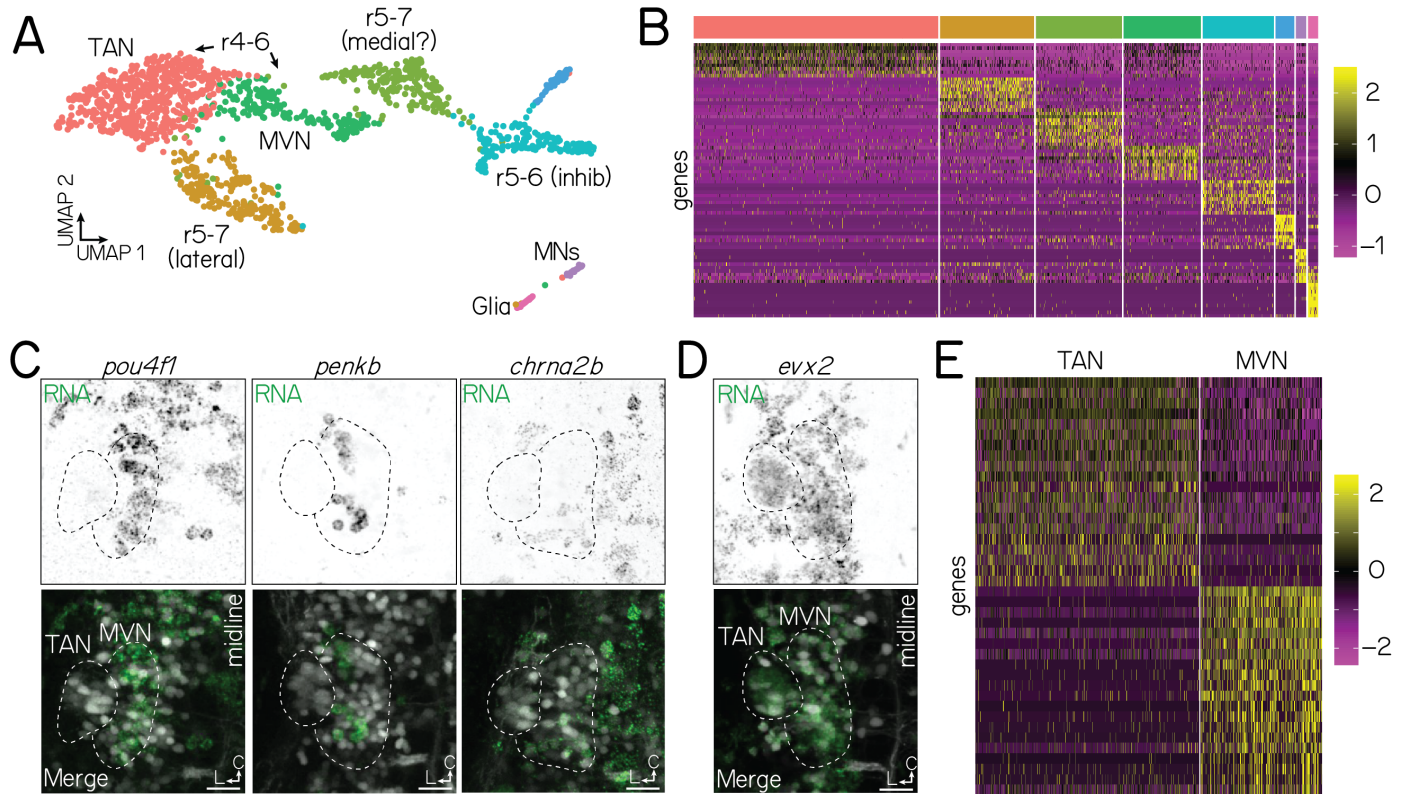


Figure S3: Molecular identification of projection neurons using a reference single-cell RNA sequencing atlas.

Associated with [Figure 5](#).

(A) UMAP visualization of a single-cell RNA sequencing atlas of $n=1,468$ neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)*, generated with 10x Genomics (Methods). Each circle is a single neuron. Neurons are clustered (colors) according to their transcriptional identity. Annotations are based on validated marker genes (data not shown). TAN, tangential nucleus; MVN, medial vestibular nucleus; r, rhombomere; MNs, motor neurons; inhib, inhibitory neurons.

(B) Heatmap showing genes unique to each annotated cluster. Each row is a gene; names unlisted for clarity. Columns show distinct clusters. Color bar on top reflects clusters in [Figure S3A](#). Yellow and purple reflect stronger or weaker gene expression, respectively.

(C) Fluorescent *in situ* hybridization against three markers (*pou4f1*, *penkb*, *chrna2b*) that are negative for tangential nucleus projection neurons and positive for medial vestibular nucleus neurons. Top row shows RNA expression (green); bottom row, merge with neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)*. Dashed lines outline the tangential nucleus (TAN) and medial vestibular nucleus (MVN). Data from 72 hpf larvae. Images shown in an axial view.

(D) Fluorescent *in situ* hybridization against a positive marker (*evx2*) for both tangential nucleus and medial vestibular nucleus neurons. All scale bars, 20 μ m.

(E) Heatmap showing genes unique to tangential and medial vestibular neurons. Clusters identified using positive and negative fluorescent *in situ* data from [Figure S3C-Figure S3D](#) and unpublished data.

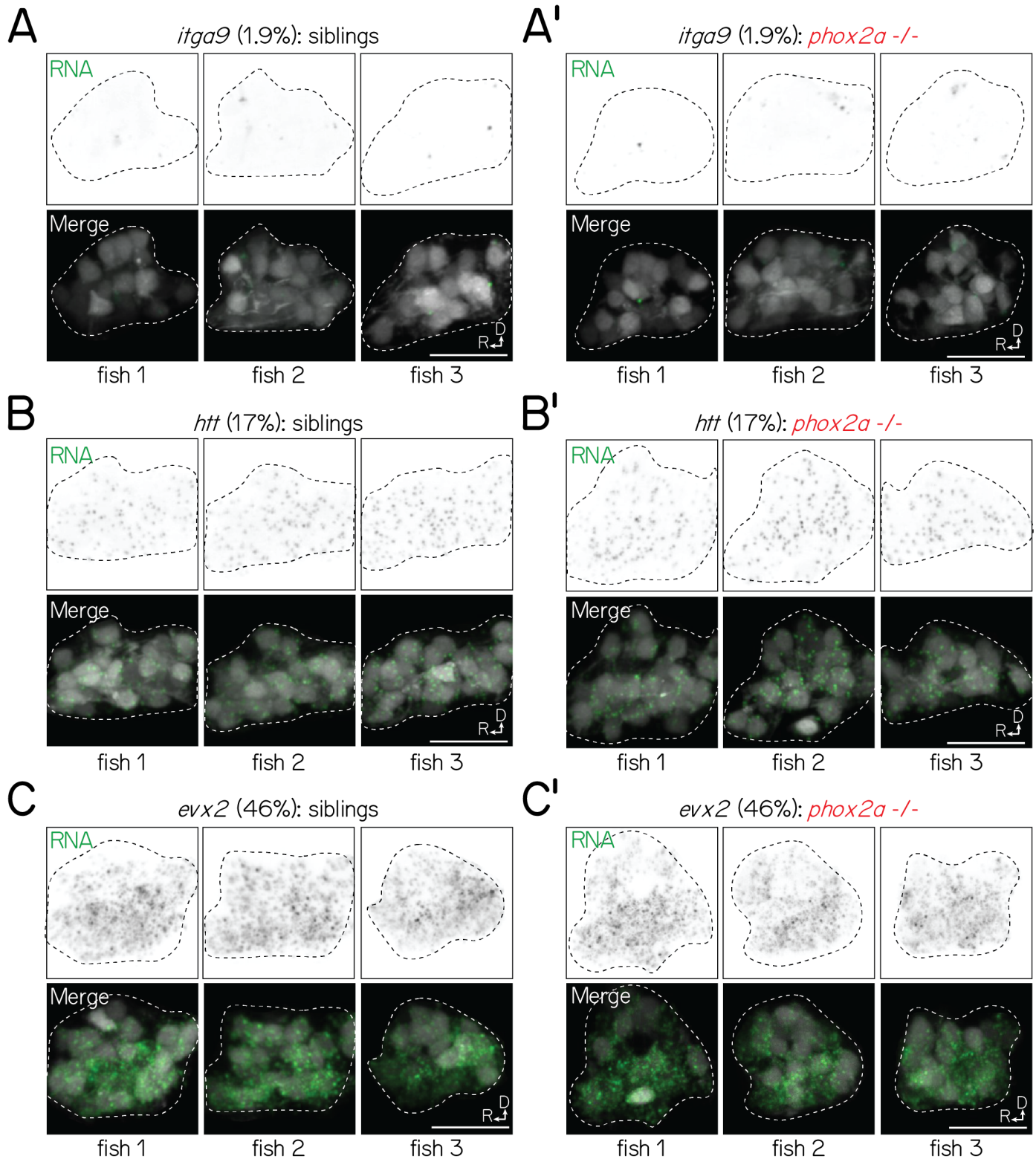


Figure S4: Visualization of transcripts in siblings and *phox2a* null mutants with fluorescent *in situ* hybridization is (1) consistent across larvae and (2) scales with predicted detection in projection neurons.

Associated with Figure 5.

(A-A') Fluorescent *in situ* hybridization against *itga9* for three sibling (A) or *phox2a* null mutant (A') larvae (72 hpf), imaged with identical conditions. Left column shows RNA (green); right column, merge with projection neurons visualized with *Tg(-6.7Tru.Hctr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (grey). Dashed lines outline the projection nucleus. Cell and transcript expression outside the projection nucleus is removed for visual clarity. Percentage (1.9%) refers to fraction of cells in a single-cell RNA sequencing reference atlas (Methods) with detected transcript. All scale bars, 20 μ m.

(B-B') Fluorescent *in situ* hybridization against *htt*, 17%, for three sibling (B) and *phox2a* mutant (B') larvae (72 hpf).

(C-C') Fluorescent *in situ* hybridization against *evx2*, 46%, for three sibling (C) and *phox2a* mutant (C') larvae (72 hpf).

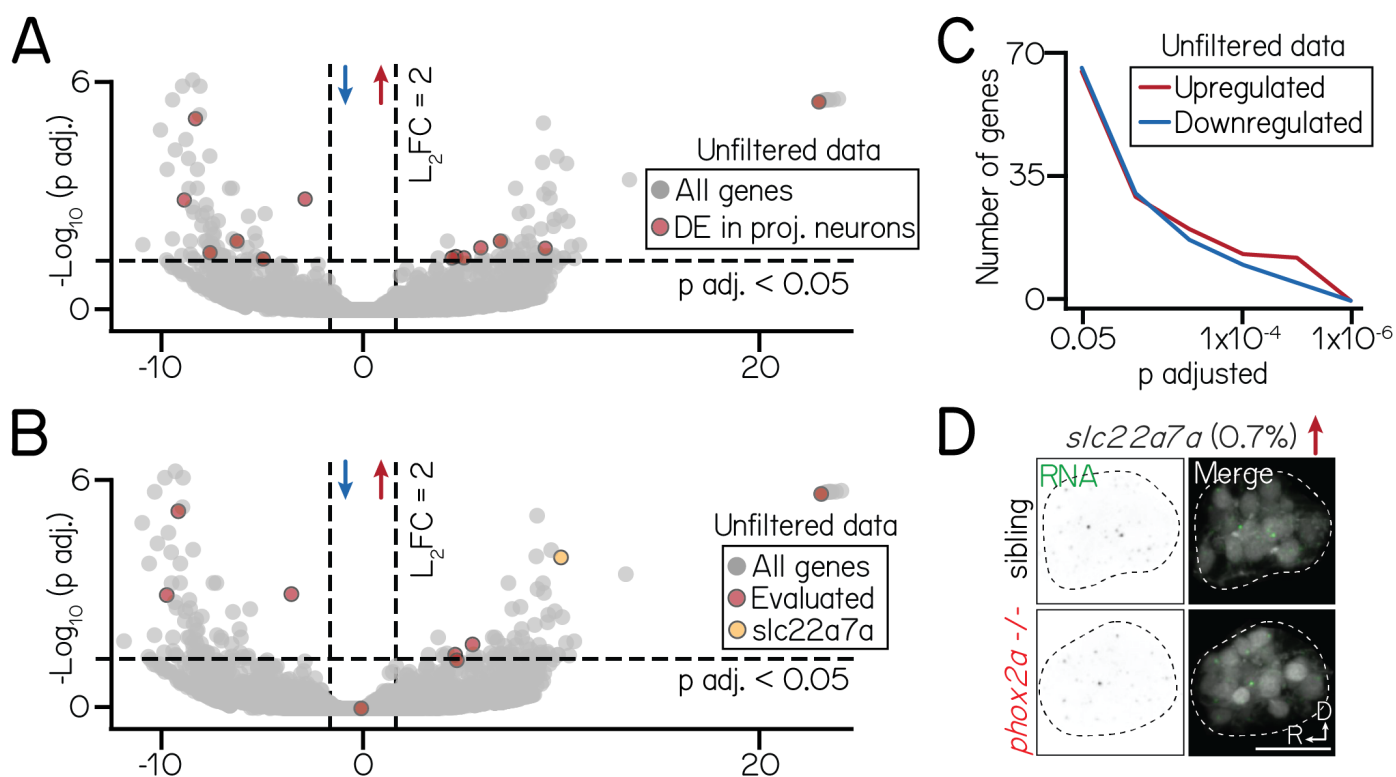


Figure S5: Differential gene expression in an unfiltered bulk sequencing dataset of siblings and *phox2a* mutants.

Associated with Figure 5.

(A) Volcano plot showing differentially expressed genes across an unfiltered bulk RNA sequencing dataset. Dashed lines represent significance cutoffs: horizontal line, p adjusted > 0.05 ; vertical line, Log_2 Fold Change > 2.0 . Each circle is a gene. Genes to the left and right of 0 on the horizontal axis show downregulated and upregulated genes, respectively. Red color shows genes that are differentially expressed in a filtered subset of projection neurons (Figure 5). Grey-colored genes are below both significance thresholds.

(B) Same data as Figure S5A, now highlighting candidate genes evaluated by fluorescent *in situ* (Figure 5) with red. One candidate (yellow) that did not meet projection neuron filter criteria (Methods) is shown in Figure S5D; remaining candidates (included in filtered data) shown in Figure 5F.

(C) Same data as Figure S5A-Figure S5B, showing the number of differentially expressed genes at progressive significance thresholds (p adjusted). Red and blue lines show the number of significantly upregulated and downregulated genes, respectively.

(D) Fluorescent *in situ* hybridization against a candidate gene, *slc22a7a* (\log_2 fold change = 10.2, p adj. = 1.6×10^{-4}), that did not meet projection neuron filter criteria. Percentage refers to fraction of projection neurons from a single-cell sequencing dataset with expression (Methods). Left columns show RNA (green); right columns, merge with projection neurons labeled with *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (grey). Dashed lines outline the projection nucleus. Cell and transcript expression outside the projection nucleus is removed for visual clarity. All scale bars, 20 μm .

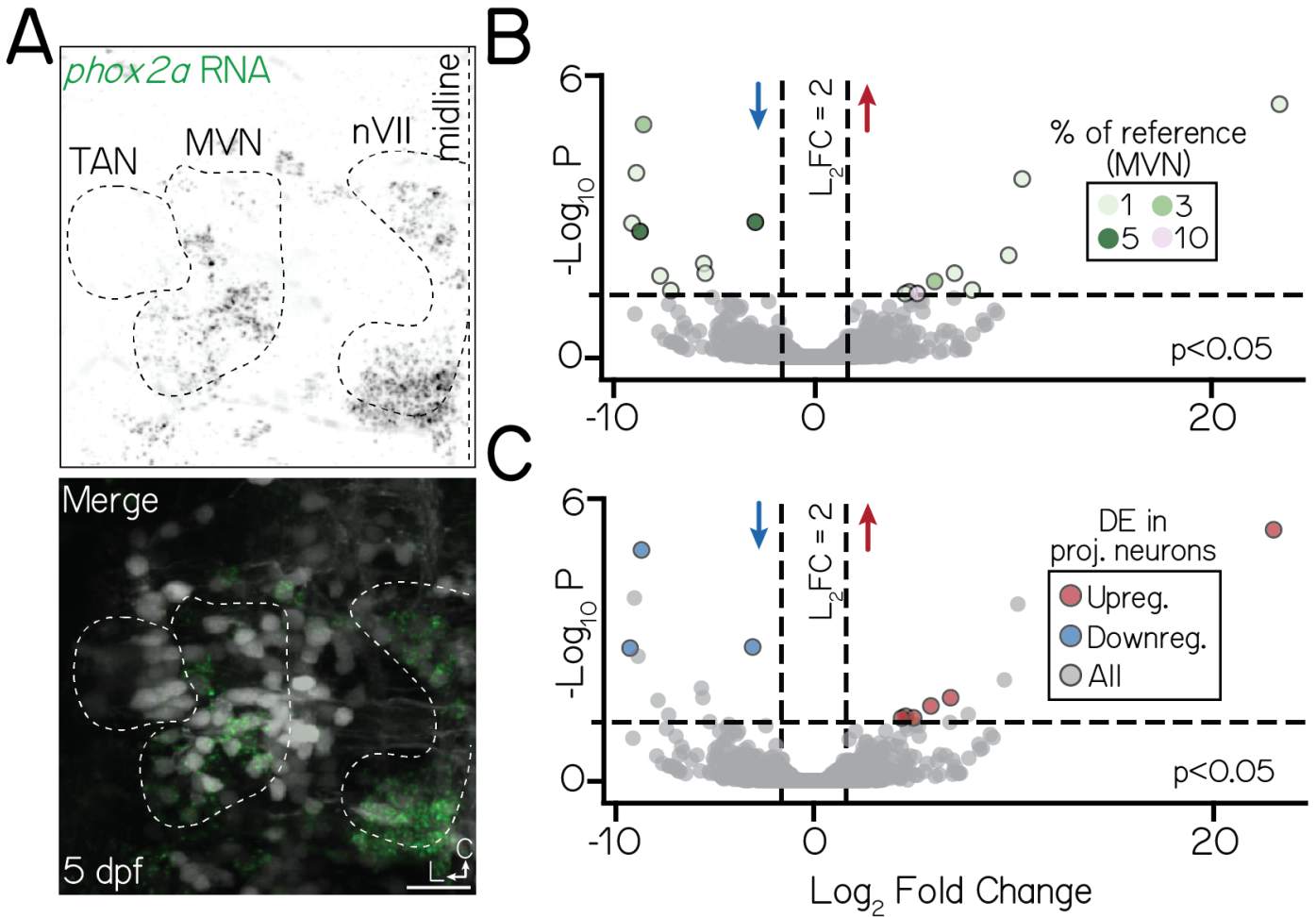


Figure S6: *phox2a* expression in the medial vestibular nucleus may underscore differential gene expression phenotypes in bulk data.

Associated with Figure 5.

(A) Fluorescent *in situ* hybridization against *phox2a* in a 5 dpf larvae (axial view). Top panel shows *phox2a* RNA (green); bottom panel, merge with neurons visualized with *Tg(isl1:GFP);Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (grey). White dashed lines outline three nuclei of interest: projection neurons in the tangential nucleus (TAN), the medial vestibular nucleus (MVN), and the facial nucleus (nVII). All scale bars, 20 μ m.

(B) Volcano plot showing differentially expressed genes in medial vestibular nucleus neurons between control and *phox2a* null larvae at 3 dpf. Dashed lines represent significance cutoffs: horizontal line, $p > 0.05$; vertical line, Log_2 Fold Change > 2.0 . Each circle is a gene. Genes to the left and right of 0 on the horizontal axis show downregulated and upregulated genes, respectively. Colors indicate percent of reference medial vestibular neurons (Methods) that express a given gene. Grey-colored genes are below both significance thresholds.

(C) Same data as Figure S6B. Color shows genes that are differentially expressed in both medial vestibular nucleus neurons and projection neurons.

	WT (all)	WT (sampled)	phox2a^{+/+}	phox2a^{+/-}	phox2a^{-/-}	p val
Tonic tilt stimuli						
n (neurons/fish)	255/10	125/x	76/5	109/6	297/16	
% sampled (nose-up/nose-down/untuned)	50/44/7	37/54/9	40/54/7	56/37/7	44/50/6	
ΔFF, nose-up	1.28±1.23	1.27±1.19	1.09±1.03	1.12±0.90	1.02±0.82	0.26
ΔFF, nose-down	2.01±1.66	1.99±1.69	1.38±0.91	1.98±1.61	2.07±1.48	0.16
directional tuning strength, nose-up	0.84±0.28	0.83±0.30	0.87±0.26	0.81±0.28	0.81±0.29	0.70
directional tuning strength, nose-down	0.72±0.30	0.72±0.31	0.73±0.31	0.77±0.30	0.68±0.29	0.54
Impulse stimuli						
n (neurons/fish)	255/10	125/x	76/5	109/6	297/16	
% sampled (responsive/unresponsive)	58/42	57/43	57/43	60/39	70/30	
ΔFF	0.41±0.46	0.33±0.28	0.29±0.29	0.22±0.16	0.32±0.28	1.0E-05
directional tuning strength	0.08±0.36	0.10±0.38	0.003±0.41	0.07±0.48	0.07±0.41	0.64
Multiple comparisons						
ΔFF to impulses	genotype		p val	Cohen's d		
	WT to sampled		p=0.13	0.21		
	WT to ^{+/+}		p=0.04	0.27		
	WT to ^{+/-}		p=3.8E-06	0.48		
	WT to ^{-/-}		p=0.006	0.24		
	^{+/+} to ^{+/-}		p=0.47	0.34		
	^{+/+} to ^{-/-}		p=0.89	0.11		
	^{+/-} to ^{-/-}		p=0.02	0.49		

Table 1: Statistical comparisons of tilt responses across genotypes. WT (sampled) refers to an n=125 neuron subset, sampled with replacement from a reference dataset of wildtype projection neurons. Data shown is mean/standard deviation unless otherwise noted. p val generated from a one-way ANOVA with multiple comparisons. Associated with [Figure 2](#) and [Figure 3](#).

Gene	% of projection neurons with expression	Putative origin	Log ₂ fold change	p adjusted
Upregulated				
* # <i>itga9</i>	1.1	r4-7	23.0	3.9E-06
# <i>dysf</i>	0.6	r4-7	6.9	0.016
<i>cers3a</i>	3.2	r4-7	9.2	0.024
* # <i>twf1b</i>	1.9	r4-7, inc. r5-6 inhib	5.9	0.024
# <i>abtb2a</i>	0.4	r4-7, inc. r5-6 inhib	4.7	0.041
* # <i>p4hb</i>	3.6	r4-7, r5-6 inhib, MNs	5.1	0.044
# <i>fhdc3</i>	1.3	r4-7, r5-6 inhib, MNs	4.5	0.044
Downregulated				
* # <i>rxfp2a</i>	2.3	r4-7, inc. inhib	-8.5	1.1E-05
* # <i>satb1a</i>	6.6	r4-7 (inc. inhib), MNs	-3.0	0.001
* # <i>bckdhbl</i>	4.2	r4-7, glia	-9.1	0.001
# <i>polrmt</i>	4.0	r4-6, inc. MVN	-8.7	0.002
CR847895.1	2.1	r4-6	-6.4	0.016
<i>asns</i>	1.5	r4-7, MNs	-7.8	0.032
BX294160.1	3.6	r4-6	-5.1	0.047

Table 2: Differentially expressed genes in projection neurons. Star indicates a gene was evaluated using fluorescent *in situ* hybridization. # symbol indicates a gene was also differentially expressed in adjacent medial vestibular neurons (see Figure S6). "% of projection neurons with expression" refers to detection in a filtered subset of projection neurons from a single-cell reference atlas of neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (Methods, Figure S3). Putative origin inferred from gene expression in the annotated 10x dataset (Methods, Figure S3)). Genes sorted by p adjusted value. Data associated with Figure 5.

Gene	% of unfiltered 10x neurons with expression	% of projection neurons
<i>ints5</i>	23.2	33.6
<i>stmn1b</i>	78.6	83.5
<i>sox4a</i>	6.5	13.1
<i>baspl</i>	61.4	56.7
<i>hmgb3a</i>	68.2	67.9
<i>ptmaa</i>	84.5	96.2
<i>gapdhs</i>	28.9	4.2
<i>pnrc2</i>	81.3	89.9
<i>snap25a</i>	65.7	46.9
<i>gpm6ab</i>	81.1	83.9
<i>calm3b</i>	0.0	0.0
<i>marcksl1b</i>	88.8	94.5
<i>tuba1c</i>	59.9	49.0
<i>cd81a</i>	43.3	36.8
<i>meis1b</i>	87.9	95.6
<i>rtn1a</i>	73.4	69.1
<i>elavl3</i>	87.1	93.4
<i>hmgb1b</i>	57.2	65.1
<i>ptmab</i>	80.7	81.2
<i>zc4h2</i>	56.9	64.3
<i>meis2b</i>	57.1	81.6
<i>slc25a5</i>	51.4	37.6
<i>mab21l2</i>	62.7	85.4
<i>h3f3c</i>	69.1	76.5
<i>rtn1b</i>	36.4	22.0
<i>elavl4</i>	78.7	83.5
<i>gng3</i>	37.2	28.3
<i>pik3r3b</i>	77.4	96.0
<i>tubb5</i>	25.3	8.9
<i>histh1l</i>	61.0	74.6
<i>serinc1</i>	51.9	45.0
<i>ckbb</i>	23.5	6.1
<i>oaz1a</i>	43.5	38.9
<i>oaz1b</i>	36.9	27.9
<i>actb1</i>	23.6	2.5
<i>ywhaba</i>	36.2	22.6
<i>ywhag2</i>	36.2	11.6
<i>si:ch211-222 21.1</i>	73.8	87.3
<i>si:dkey-276j7.1</i>	45.8	41.4
<i>aldocb</i>	19.3	2.5
<i>actb2</i>	27.0	5.1
<i>tmem59l</i>	39.8	24.5
<i>calm2b</i>	37.9	19.5
<i>hmgn6</i>	73.6	87.9
<i>h2afx1</i>	59.6	64.7
<i>cd99l2</i>	32.4	20.3
<i>cirbpb</i>	77.8	88.2
<i>ppdpfb</i>	74.5	90.5
<i>stxbp1a</i>	52.3	32.6
Control		
<i>evx2</i>	33.8	50.0

Table 3: Top 50 expressed genes in an unfiltered bulk RNA sequencing dataset of *phox2a* siblings. "% of unfiltered 10x neurons" refers to gene detection in a single-cell atlas of neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (n=1,468 neurons). "% of projection neurons" refers to gene detection in a subset of the single-cell atlas containing projection neurons in the tangential nucleus (n=473 neurons). Data associated with [Figure 5](#).

Gene	% of unfiltered 10x neurons with expression	Putative origin	Log ₂ fold change	p adjusted
Upregulated				
<i>macc1</i>	0.1	r4-6	24.0	3.2E-06
<i>CR55941.1</i>	0.0		23.7	3.4E-06
<i>si:dkey-65b12.6</i>	0.0		23.5	3.4E-06
<i>si:ch73-106n3.2</i>	0.1		23.5	3.4E-06
<i>mcm10</i>	0.1	MNs	23.4	3.4E-06
<i>si:ch211-244o22.2</i>	0.5	r4-6	23.4	3.4E-06
<i>dre-mir-10a</i>	0.0		23.3	3.5E-06
<i>itga4</i>	0.3	r5-6 (inhibitory)	23.2	3.5E-06
<i>si:dkeyp-87d8.8</i>	0.0		23.2	3.6E-06
<i>arsj</i>	0.5	MNs	23.0	3.9E-06
<i>tlr1</i>	0.0		23.0	3.9E-06
*% <i>itga9</i>	2.3	r4-7	23.0	3.9E-06
<i>otofb</i>	0.5	r4-6	9.1	1.4E-05
<i>myo7ba</i>	0.4	r4-7	9.8	1.0E-04
<i>zfand1</i>	0.1	MNs	9.0	1.5E-04
% <i>slc22a7a</i>	0.7	r4-7	10.2	1.5E-04
<i>agrp</i>	0.0		13.4	4.1E-04
<i>si:dkey-46i9.6</i>	0.1	r5-7	7.7	6.8E-04
<i>muc2.2</i>	0.0		9.4	6.9E-04
<i>cd37</i>	0.0		9.1	9.8E-04
<i>musk</i>	0.3	r4-6	9.4	1.2E-03
<i>mcamb</i>	0.2	r5-7	8.3	2.7E-03
<i>ppp1r42</i>	0.5	r5-6 (inhibitory)	7.9	3.1E-03
<i>CR677513.1</i>	0.0		9.9	3.5E-03
Downregulated				
*% <i>satb1a</i>	7.9	r4-7 (inc. inhib), MNs	-8.6	1.0E-06
<i>znf975</i>	0.7	r4-6	-8.3	1.5E-06
<i>phldb1a</i>	0.6	r5-7 (inc. inhib)	-9.2	1.5E-06
<i>TSTA3</i>	0.0		-9.7	3.4E-06
<i>si:dkey-24p1.6</i>	0.0		-8.3	8.4E-06
<i>si:dkey-77f5.14</i>	0.2	r5-7	-8.5	1.1E-05
<i>tha1</i>	0.1	MVN	-10.3	2.1E-05
<i>serpinh2</i>	0.5	r4-6	-9.0	3.7E-05
<i>ghrh</i>	0.3	r4-7	-9.5	6.9E-05
<i>asah1b</i>	0.8	r4-7	-7.8	9.9E-05
<i>msmo1</i>	0.9	r5-7, inc. inhib	-8.9	1.1E-04
<i>tagln2</i>	0.3	glia	-8.4	2.2E-04
<i>zgc:174863</i>	0.1	MNs	-9.9	2.2E-04
*% <i>rxfp2a</i>	3.2	r4-7, inc. inhib	-6.6	6.8E-04
<i>bmp4</i>	0.7	r4-7	-6.8	6.8E-04
<i>cfl1l</i>	0.1	r4-6	-8.4	6.8E-04
*% <i>polrmt</i>	4.2	r4-7, inc. inhib	-8.8	6.9E-04
<i>anxa2a</i>	0.6	r4-7	-3.0	1.3E-03
<i>galr1a</i>	0.3	MVN	-9.1	1.4E-03
<i>selenow2b</i>	0.1		-8.0	1.8E-03
*% <i>bckdhbl</i>	1.4	r4-7, glia, MNs	-8.7	2.1E-03
<i>boka</i>	0.5	r5-7	-8.6	2.9E-03
<i>cyldb</i>	0.2	r4-7	-7.9	3.0E-03
<i>pon2</i>	0.6	r4-7, glia, MNs	-5.0	3.1E-03
<i>si:ch73-204p21.2</i>	0.3	r5-7, inc. inhib	-8.2	3.5E-03
<i>and2</i>	0.1	r4-6	-5.8	3.7E-03
Control				
<i>evx2</i>	33.8	r4-7	0.46	0.99

Table 4: Top 50 differentially expressed genes in an unfiltered bulk RNA sequencing dataset of *phox2a* siblings and null mutants. One star indicates a gene was retained in a filtered subset of projection neurons; %, evaluated using fluorescent *in situ* hybridization. "% of unfiltered 10x neurons" refers to gene detection in an unfiltered single-cell reference atlas of neurons labeled in *Tg(-6.7Tru.Hcrtr2:GAL4-VP16);Tg(UAS-E1b:Kaede)* (n=1,468 neurons). Putative origin inferred from gene expression in the annotated 10x dataset (Methods, Figure S3)). Genes sorted by p adjusted value. Data associated with Figure 5.

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, peptides, and recombinant proteins		
Tween	Fisher Scientific	BP337-100
32% paraformaldehyde	Electron Microscopy Sciences	15714
Proteinase K	ThermoFisher Scientific	25530049
Papain	Worthington Biochemical	LK003178
Hanks Buffered Salt Solution (HBSS)	ThermoFisher Scientific	14170112
Earl's Buffered Salt Solution (EBSS)	ThermoFisher Scientific	24010043
DNAse	Worthington Biochemical	LK003172
DAPI	Invitrogen	D1306
L15 Medium	ThermoFisher Scientific	11415064
Fetal bovine serum, qualified, triple-filtered	ThermoFisher Scientific	A3160501
Collagenase Type 1A	Sigma Aldrich	C9891-500MG
Low melting point agarose	ThermoFisher Scientific	16520
Ethyl-3-aminobenzoic acid ethyl ester (MESAB)	Sigma Aldrich	E10521
Pancuronium bromide	Sigma Aldrich	P1918
Critical commercial assays		
<i>in situ</i> hybridization chain reaction v3.0 (HCR)	Molecular Instruments	N/A
RNAqueous Micro Total RNA Isolation Kit	ThermoFisher Scientific	AM1931
MEGashortscript T7 Transcription Kit	ThermoFisher Scientific	AM1354
QiaQUICK PCR Purification Kit	Qiagen	28104
EnGen Spy Cas9 NLS	New England Biolabs	M0646T
Deposited data		
Raw and analyzed calcium imaging data	This study	DOI: 10.17605/OSF.IO/93V6E
Raw and analyzed 10x Genomics scRNA-seq datasets	This study	GEO: GSE254346
Raw and analyzed bulk RNA seq datasets	This study	GEO: GSE254345
Experimental models: Organisms/strains		
<i>Tg(-6.7Tru.Hcrtr2:GAL4-VP16)</i>	25,49	ZFIN: ZDB-TGCONSTRUCT-151028-8
<i>Tg(UAS-E1b:Kaede)</i>	48	ZFIN: ZDB-TGCONSTRUCT-070314-1
<i>Tg(isl1:GFP)</i>	40	ZFIN: ZDB-ALT-030919-2
<i>Tg(UAS:GCaMP6s)</i>	42	ZFIN: ZDB-TGCONSTRUCT-140811-3
<i>phox2a^{d22}</i>	This study	N/A
<i>phox2a^{d19}</i>	This study	N/A
<i>phox2aⁱ²</i>	This study	N/A
Oligonucleotides		
<i>phox2a</i> forward primer (5-CAGCCAGAGCAACGGCTTCC-3)	Sigma Aldrich	N/A
<i>phox2a</i> reverse primer (5-AAGCCGACAACAGTGTGTGTAA-3)	Sigma Aldrich	N/A
<i>phox2a</i> guide 1 (5-CTCGCCACCGCCAGCTGCAC-3)	Sigma Aldrich	N/A
<i>phox2a</i> guide 2 (5-CTCGGCTTCAGCTCCGGCC-3)	Sigma Aldrich	N/A
HCR probes	Integrated DNA Technologies	N/A
Software and algorithms		
Fiji/ImageJ	124	RRID: SCR_02285
Adobe Illustrator (2021)	Adobe	RRID: SCR_010279
Matlab 2020b	Mathworks	RRID: SCR_001622
Seurat v4	133	https://satijalab.org/seurat
CRISPR Guide RNA Design Tool	Benchling	https://benchling.com/crispr
Other		
20 micron cell strainer	pluriSelect	431002060
SH800z 100 micron sorting chip	Sony	LE-C3210

Table 5: Key Resources Table (associated with Methods).