NEUROSCIENCE

A single-cell resolution gene expression atlas of the larval zebrafish brain

Inbal Shainer⁺, Enrico Kuehn⁺, Eva Laurell, Mariam Al Kassar, Nouwar Mokayes, Shachar Sherman, Johannes Larsch, Michael Kunst[‡], Herwig Baier^{*}

The advent of multimodal brain atlases promises to accelerate progress in neuroscience by allowing in silico queries of neuron morphology, connectivity, and gene expression. We used multiplexed fluorescent in situ RNA hybridization chain reaction (HCR) technology to generate expression maps across the larval zebrafish brain for a growing set of marker genes. The data were registered to the Max Planck Zebrafish Brain (mapzebrain) atlas, thus allowing covisualization of gene expression, single-neuron tracings, and expertly curated anatomical segmentations. Using post hoc HCR labeling of the immediate early gene *cfos*, we mapped responses to prey stimuli and food ingestion across the brain of freely swimming larvae. This unbiased approach revealed, in addition to previously described visual and motor areas, a cluster of neurons in the secondary gustatory nucleus, which express the marker *calb2a*, as well as a specific neuropeptide Y receptor, and project to the hypothalamus. This discovery exemplifies the power of this new atlas resource for zebrafish neurobiology.

INTRODUCTION

Development and function of neural circuits depend on the spatiotemporal dynamics of differential gene expression across the brain (1, 2). Particular classes of genes play prominent roles in organizing the layout of the nervous system and may serve as markers of celltype identity and/or function. These include genes whose protein products directly influence cell-cell interactions and local properties of the synaptic network and those that regulate the expression of other genes to determine neuronal cell fate, diversification, morphology, and connectivity (3-6). Often the corresponding RNAs are expressed in a region- and cell type-specific fashion. Another class of important marker genes encode functional determinants, e.g., neurotransmitter-synthesizing enzymes, ion channels, synaptic machinery components, and neuropeptides; the presence of the corresponding proteins provides information about physiological properties of the expressing neurons and tends to be not cell-type specific (except for neuropeptides). Visualizing the expression of these markers in an atlas format offers a glimpse into the cellular constituents of neural circuitry and their molecular endowment (5, 6). Such an atlas then allows researchers to select markers for labeling of neurons of interest and genetic handles for their manipulation. Moreover, covisualizing many of these markers offers a holistic view of the genetic architecture of the brain and may reveal general organizing principles.

The most prominent attempt at visualizing brain-wide transcript expression, while preserving spatial layout of the tissue, is probably the Allen Brain Atlas project, which published the results of RNA in situ hybridizations (ISHs) for thousands of genes in tissue sections, first in mouse (5) and later in human (6). These pioneering studies, together with both prior and more recent work with similar aims

*Present address: Allen Institute for Brain Science, Seattle, WA 98109, USA.*Corresponding author. Email: herwig.baier@bi.mpg.de

[‡]Present address: Allen Institute for Brain Science, Seattle, WA 98109, USA.

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[e.g., (7–9)], are of immense value for the research community. The resulting resources are routinely consulted by experimenters seeking to corroborate hypotheses and identify genetic markers for neuronal subpopulations of interest.

Atlases of the nervous system have also been devised for other model organisms, although, to our knowledge, not yet for RNA ISH patterns to faithfully map the endogenous gene expression. The nervous system of the nematode worm Caenorhabditis elegans has been charted in unsurpassed detail (10). Most recently, a strain of C. elegans has been developed by genetic engineering in which each of its 302 neurons is identifiable in one animal by a combination of fluorescent proteins (11). This multicolor atlas enables in vivo interrogation of the spatial arrangement of identifiable neurons and serves as a convenient readout of the effects of developmental perturbations on cell fate and cell positioning (11). In the fruitfly Drosophila melanogaster, the "virtual fly brain browser" (12) and the "fruit fly brain observatory" (13) offer standardized neuroanatomical frames in which thousands of transgene expression patterns are coregistered, generating a light-microscopic mesoscale connectome.

In the zebrafish Danio rerio, histological brain atlases have been available for some time [adult: (14, 15); embryo/larva: (16)], as well as atlas databases of hundreds of transgenic lines and immunostainings, such as "ViBE-Z" (17), "Z-brain" (18), and the "Zebrafish Brain Browser" (19). The Max Planck Zebrafish Brain ("mapzebrain") atlas (20) at https://mapzebrain.org is a multimodal digital atlas of the 6-day postfertilization (dpf) larval brain, integrating data portals for transgenic lines, histological stains, and expertly curated neuroanatomical regions, as well as a growing number (currently ca. 4300) of single-neuron morphologies. These datasets are all registered into the same spatial coordinate system, the so-called standard brain (20). The mapzebrain atlas has also been cross-linked to Zbrain and the Zebrafish Brain Browser to enable data transfer between the different resources (20). Its Web interface offers refined visualization and animation tools, as well as options for downloading original or processed data for offline analyses.

Max Planck Institute for Biological Intelligence, Martinsried, Germany. †These authors contributed equally to this work.

Here, we set out to add RNA fluorescent in situ hybridization (FISH) patterns to the mapzebrain atlas. Single-cell resolution was achieved by hybridization chain reaction (HCR), a spatial transcriptomic method that is highly sensitive, specific, and multiplexable, allowing visualization of several RNA species in one specimen with different probe-specific fluorophores (*21*). Moreover, as a powerful tool for recording activity hotspots across the brain, we have devised an HCR FISH protocol for detecting *cfos* RNA following various sensory stimulations and behavioral tasks. All the data are publicly available and free to download through the mapzebrain website (https://mapzebrain.org/). This new resource, which can be straightforwardly expanded in the future by community contributions, offers convenient Web-based access to markers of neuronal subpopulations and paves the way for a genetic analysis of circuit development and structure in this widely used vertebrate model.

RESULTS

Pipeline for HCR in situ labeling and registration to the mapzebrain atlas

Recent developments of FISH techniques, such as RNAscope, MERFISH, and HCR (9, 21, 22), offer single RNA molecule detection and multiplexing capabilities. We chose HCR, because the staining protocol is easily implemented in larval and juvenile fish (21, 23) and is suitable for registration into a common three-dimensional (3D) reference (23, 24). HCR technology enables multiplexed FISH of up to five probe sets per round of staining with different fluorophores (21). To scale up the number of RNA species mapped, we tested two approaches: either multiround HCR FISH by stripping the HCR signal by deoxyribonuclease I treatment and restaining of the same sample (24, 25) or labeling of different genes in different specimens and computational registering into a common coordinate system. Although the former strategy gives theoretically unlimited information about coexpression in individual cells, we decided against it mainly for two reasons: First, we noticed that iterative rounds of stripping and relabeling, together with repeated agarose embedding for imaging, lead to a reduced fluorescence of the reference channel [*Tg(elavl3:H2b-GCaMP6s)*; see below] and a weaker in situ staining. Second, the initial set of markers is, of course, not final, as new markers or better probes are certain to become available in the future. Thus, to allow for seamless addition of new data to the atlas, we chose to generate expression maps from single rounds of HCR and computational coregistration into a standard brain.

To enable coalignment of different brains, we carried out HCR FISH stainings in the background of the *Tg(elavl3:H2b-GCaMP6s)* transgenic line, in which nuclear-localized GCaMP6s is expressed in almost all neurons. We generated an averaged brain template of the *Tg(elavl3:H2b-GCaMP6s)* line using the Advanced Normalization Tools (ANTs) toolkit (fig. S1) (*26*) and calibrated a registration protocol (see Methods; movie S1). We then acquired FISH data for 290 genes (table S1), which were previously identified as distinct, often cell type–specific, markers in retina, tectum, and diencephalon (*27–29*). This initial set of markers includes genes encoding transcription factors, cell surface and secreted molecules, enzymes involved in transmitter synthesis and transport, neuropeptides, calcium-binding proteins, and other factors (Fig. 1 and table S1).

For each marker gene, we labeled and imaged three 6-dpf fish, which we registered to the standard brain and averaged to ensure

we are mapping a consistent expression signal among animals (Fig. 1; see Methods). We multiplexed two genes per sample (table S1) and acquired the confocal data at single-cell resolution, revealing the number of cells expressing a particular gene per region of interest (ROI), as well as any overlapping expression (Fig. 2). The confocal images were acquired from dorsal to ventral; this resulted in a reduced signal in the deep ventral parts due to light scattering. We therefore implemented a signal enhancement visualization option at mapzebrain.org to better explore the ventral expression patterns. All the expression patterns of individual marker genes, including those in individual and averaged brains, are publicly accessible through the latest version of mapzebrain at http://mapzebrain.org/ (Fig. 1).

FISH image coregistration with other mapzebrain data modalities

Coregistration of the new FISH dataset with existing mapzebrain datasets frequently revealed spatial overlaps. For example, after registration, the expression of *aldh1a2* partially overlapped with the pattern of the transgenic gene trap line *mpn321Gt* in the pretectum (fig. S2). To verify this in silico overlap of registered data acquired by different procedures (i.e., live imaging, immunostaining, and FISH), we carried out *aldh1a2* FISH labeling in larvae expressing the *mpn321Gt* transgene and indeed found coexpression of the two markers in individual cells in the pretectum (fig. S2). This finding highlights the potential to identify and confirm marker genes for cell populations of interest via rapid FISH labeling and demonstrates the robustness and reproducibility of our digital image registration pipeline across samples and data acquisition procedures.

Generating post hoc cfos brain activity maps of freely swimming fish

To expand the capabilities of our gene expression resource, we challenged the fish larvae in sensory or behavioral paradigms, for a total of 40 min, and carried out post hoc FISH staining of RNA from the immediate early gene *cfos* (Fig. 3, fig. S3, and movie S2). The *cfos* gene, also known as *fosab*, is expressed in neurons after prolonged depolarization and is commonly used as a marker integrating recent neuronal activity (*30*, *31*).

We exposed two groups of larval zebrafish to high temperature (32°C) or low temperature (22°C). Another group was allowed to hunt prey. Other larvae were held on a black, or a white, background. Still, others were subjected to a drifting, black-and-white grating, which induced an optomotor response (32, 33), or to a repeatedly looming black disk on a white background, thus evoking a series of visual escape responses (34). Two groups were exposed either to an asymmetric light source, which evoked phototaxis (35, 36), or to the alarm substance, "Schreckstoff," which was extracted from fish skin (see Methods). For each condition, we imaged, registered, and analyzed six larvae. Qualitative inspection of the cfos FISH-labeled areas revealed specific activity hotspots for different behavioral conditions (Fig. 3, fig. S3, and movie S2). Such activity maps not only largely confirmed previously published calcium imaging studies of embedded animals (34, 37-40) but also revealed previously unidentified hotspots. For example, in larvae that were hunting prey, we observed strong cfos labeling in the anterior optic tectum and retinal arborization field 7 (AF7; Fig. 3, fig. S3, and movie S2), which have been implicated in prey detection



Fig. 1. Gene expression atlas of the larval zebrafish brain: Pipeline and example patterns. (**A**) Multiplexed FISH was performed on 6-dpf *Tg(elav13:H2b-GCaMP6s)* fish. The GCaMP6s signal was used to align the images to the reference brain, and the aligned images were then uploaded to the mapzebrain atlas. Scale bar, 100 μm. (**B**) 3D projections of 50 selected FISH images, registered into the standard brain. Color coding: green, enzymes; cyan, transcription factors; red, neurotransmitter synthesizing enzyme (GABAergic) and transmitter transporters (glutamatergic); magenta, neuropeptides; yellow, miscellaneous other. Scale bar, 100 μm.

and prey capture initiation in both embedded and freely swimming fish (40-44), as well as in cells of the secondary gustatory nucleus (SGN), which is located adjacent to the cerebellar commissure (Fig. 3 and movie S2) (45-47).

Molecular and cellular analysis of a brain area associated with hunting

Since activity in the SGN had not been reported in brain-wide imaging studies in embedded animals, we suspect that it is contingent on the animal to freely swim and hunt. Since it is likely driven by gustatory inputs, we propose that its activation is linked to the ingestion of food rather than the detection of prey or huntingrelated movement (42, 44). To further explore how *cfos* maps can be used for circuit analysis, we interrogated the mapzebrain atlas for marker genes overlapping with the *cfos* signal and found that this region contains a group of neurons labeled by *calb2a* (*calreti-nin*), a known marker for the SGN (46) and its corresponding anti-Calretinin immunostaining. We verified coexpression of *cfos* and *calb2a* by multiplexed HCR FISH (Fig. 4, A to C).



Fig. 2. Multiplexed HCR in situ labeling of single cells. Example of a single focal plane of a multiplexed HCR labeled fish. The endogenous expression of *Tg(e-lavl3:H2b-GCaMP6s)* labels neuronal nuclei and can be used to segment single neurons and to identify whether genes (here *cart2* and *cart3*) are coexpressed (top right panel, arrowheads) or not (bottom right panel). Scale bars, 100 μ m (left) and 10 μ m (right).

We scanned other gene expression maps and discovered that *npy1r*, which encodes a neuropeptide Y (NPY) receptor, also overlaps with the SGN-associated *cfos* pattern (Fig. 4D). NPY has been implicated in the regulation of food consumption and foraging (*48–50*), and the expression map of *npy1r* suggests that NPY could function as a modulator of the gustatory pathway through direct signaling to SGN neurons. Other NPY receptors that were examined did not express within the SGN area (Fig. 4D) but showed various, complementary expression patterns in other brain areas. Nonoverlapping expression of the NPY receptor repertoire in distinct neuronal subpopulations may underlie local specificity of modulation by this neuropeptide.

Next, we drew a 3D ROI encompassing the *cfos*-labeled region near the SGN to query the database for individual, morphologically reconstructed neurons (see movie S3 for a detailed tutorial). We used the search feature of mapzebrain and identified 32 neurons that had their somata within the 3D ROI (Fig. 4E). SGN neurons were hierarchically clustered into four morphological types using the NBLAST algorithm (51). One of these clusters is located at the edge of the 3D ROI mask, evidently in the cerebellum, and is composed of Purkinje cells (cluster 2; Fig. 4E). Other groups project to the contralateral hindbrain (cluster 1) or to the tectum and the contralateral AF7 through the postoptic commissure (cluster 3). Notably, one cluster of SGN neurons (cluster 4) extends axons to the dorsal zone of the lateral hypothalamus (Fig. 4E), an area that is activated in response to both prey stimuli and satiety (39, 52) and which also shows strong cfos expression following hunting behavior (movie S2). Thus, exploring cfos activity maps with morphologically reconstructed neurons revealed connections between SGN, AF7-associated pretectum, tectum, and hindbrain, brain areas that are jointly engaged during pursuit and consummation of prey.

DISCUSSION

Here, we present an essential addition to the neuroscience resources for larval zebrafish: whole-brain, single-cell resolution gene expression maps for an initial set of 290 markers, coregistered with thousands of single-neuron tracings, hundreds of transgenic lines, and more than 120 expertly curated anatomical segmentations in a common atlas space. For more than 20 years, the main zebrafish RNA expression resource has been the one produced by Thisse and Thisse (53, 54). It provides colorimetric whole-mount ISH images in 2D, which are accessible through the Zebrafish Information Network (54, 55). While this resource contributed greatly to zebrafish research in many fields, its utility for neuroscience is limited: Most of the data were collected for early developmental stages (up to 72 hours postfertilization), do not offer cellular resolution or spatial accuracy due to the colorimetric ISH method used,



Fig. 3. *cfos* **brain activity maps of freely swimming fish.** 3D projections of *cfos* FISH-labeled areas of neuronal activity during behavior. Dorsal views are shown for each experimental condition. For interactive viewing, spanning the full depth of the brain, see mapzebrain atlas at https://mapzebrain.org/. Scale bar, 100 μm.

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Fig. 4. Marker gene expression and hypothetical connectivity of SGN neurons involved in prey capture. (**A**) Orthogonal views of the *cfos* prey capture brain activity map with the registered *calb2a* FISH and anti-Calretinin immunostaining. An ROI is drawn around the *cfos*-positive SGN area (yellow). Scale bar, 100 μm. A, anterior; D, dorsal; P, posterior; V, ventral. (**B**) Enlargement of the *cfos*-positive SGN area. Possible colabeling was identified between the registered *cfos* and *calb2a* data. (**C**) Multiplexed FISH of *calb2a* and *cfos* verified the suggested colabeling (arrowheads). A single focal plane of an individual larva is shown. Scale bar, 10 μm. (**D**) Expression patterns of several NPY receptors in the same single *z* plane (z261). The yellow line labels the *cfos* + SGN area. *npy1r* shows strong expression within the SGN, which is not seen for the other tested NPY receptors. Scale bar, 100 μm. (**E**) The ROI drawn in (A) was used to search for single neurons whose somas are within the ROI. Thirty-two neurons were identified, mirrored onto the left hemisphere [see (20)] and hierarchically clustered according to their morphology. One cluster of Purkinje cells (cluster 2) projecting to the octavolateralis nucleus was identified, as well as three clusters of SGN neurons, with projections to the contralateral hindbrain (cluster 1), to the contralateral retinal AF7 and tectum through the postoptic commissure (cluster 3), and to the hypothalamus (cluster 4).

and cannot be registered to the newly emerging zebrafish brain atlases. Our new dataset closes this gap.

As an added feature, we provide *cfos* brain activity maps for several commonly studied behaviors of larval zebrafish. The *cfos* method has at least two experimental advantages over other post hoc activity labeling methods: (i) *cfos* expression is localized to the soma, without neuropil labeling, which improves cell detection and segmentation compared to phosphorylated extracellular-signal related kinase (pERK) detection (*14*) and CAMPARI (*52*). (ii) *cfos* labeling can be combined with cell marker probes by iterative (*20*) or multiplexed HCR stains.

In proof-of-concept experiments, we have succeeded in linking *cfos* maps with other data modalities, such as gene expression and single-cell connectivity. This has enabled the discovery of a behaviorally relevant pathway. The *calb2a*⁺ SGN neurons are activated during hunting by gustatory cues (Fig. 4) (*38*, *40*). By using the atlas query tool (*20*) and gene expression maps, we identified their projection patterns, which likely convey this information to an area in the lateral hypothalamus (*39*). In addition, SGN neurons also express *npy1r* and are thus targets of modulation by the orexigenic peptide NPY. While an experimental test of these hypotheses is beyond the scope of the current work, these findings illustrate the power of integrating gene expression, functional *cfos* labeling, transgene, and single-cell reconstructions in a multimodal digital atlas.

The Web browser interface of the mapzebrain atlas offers advanced computational, visualization, and downloading options. Bridging registrations between mapzebrain and other atlases have been made available (20). Moreover, standardized protocols for staining and imaging enable coregistration of data acquired in different laboratories, supporting different microscopes and imaging modalities (56, 57). The mapzebrain atlas thus offers a platform for covisualization of gene expression and cellular architecture and a starting point for explorations of circuit function and structure. Future work will be directed at integrating brain-wide functional imaging data (e.g., from calcium or voltage recordings) and electron microscopic reconstructions (58) into the atlas.

METHODS

Zebrafish husbandry and maintenance

Adult zebrafish of the TLN strain were kept at 28° C with a day/night cycle of 14/10 hours, pH 7 to 7.5, and a conductivity of 600 µS. All the experiments were performed on *Tg(elavl3:H2b-GCaMP6s)* larvae produced by natural matings. Larvae were raised until 6 dpf at 28° C in Danieau's solution in petri dishes. The larvae were treated with phenylthiocarbamide (phenylthiourea, PTU) starting 24 hours postfertilization to prevent pigmentation. For *cfos* experiments, *Tg(elavl3:H2b-GCaMP6s)* larvae in the TLN background, carrying the *mitfa* mutation and therefore lacking melanin pigment, were used without PTU treatment. The animal experiments were performed under the regulations of the Max Planck Society and the regional government of Upper Bavaria (Regierung von Oberbayern) approved protocols: ROB-55.2-2532.Vet_02-20-11, ROB-55.2-2532.Vet_02-19-16.

HCR FISH staining

HCR reagents, including probes, hairpins, and buffers, were purchased from Molecular Instruments (Los Angeles, CA, USA). The staining was performed according to a modified protocol of the "HCR RNA-FISH protocol for whole-mount zebrafish embryos and larvae" provided by Molecular Instruments (21). Tg(elavl3:H2b-GCaMP6s)-positive larvae were anesthetized in 1.5 mM tricaine and fixed with ice-cold 4% paraformaldehyde (PFA)/Dulbecco's phosphate-buffered saline (DPBS) overnight at 4°C with gentle shaking. The following day, larvae were washed three times for 5 min with DPBST ($1 \times$ DPBS + 0.1% Tween 20) to stop fixation, followed by a short 10-min treatment with ice-cold 100% methanol (MeOH) at -20°C to dehydrate and permeabilize the tissue samples. Next, rehydration was performed by serial washing of 50% MeOH/50% DPBST and 25% MeOH/75% DPBST for 5 min each and lastly 5×5 min in DPBST. Ten to 12 larvae were transferred into a 1.5-ml Eppendorf tube and prehybridized with prewarmed hybridization buffer for 30 min at 37°C. Probe solution was prepared by transferring 2 pmol of each HCR probe set (2 µl of 1 µM stock) to 500 µl of hybridization buffer at 37°C. For cfos experiments, 4 µl of 1 µM probe stock was used, while for genes with relatively weak expression levels (i.e., receptors), 8 μ l of 1 μ M probe stock was used. The hybridization buffer was replaced with probe solution, and the samples were incubated for 12 to 16 hours at 37°C with gentle shaking. To remove excess probe, larvae were washed 4×15 min with 500 µl of prewarmed probe wash buffer at 37°C. Subsequently, larvae were washed 2×5 min with $5 \times$ SSCT ($5 \times$ sodium chloride sodium citrate + 0.1% Tween 20) buffer at room temperature. Next, preamplification was performed by incubating the samples in 500 µl of amplification buffer for 30 min at room temperature. Separately, 30 pmol of hairpin h1 and 30 pmol of hairpin h2 were prepared by snap-cooling 10 µl of 3 µM stock by incubating the hairpins in 95°C for 90 s and cooling down to room temperature in a dark environment. After cooling down for 30 min, hairpin solution was prepared by transferring the h1 and h2 hairpins to 500 µl of amplification buffer. The preamplification buffer was removed, and the samples were incubated in the hairpin solution for 12 to 16 hours in the dark at room temperature. Excess hairpins were washed the next day 3×20 min using $5 \times$ SSCT at room temperature. Larvae were then long-term stored at 4°C in 5× SSCT until imaging.

Imaging

Samples were embedded in 2% low-melting agarose in 1× DPBS and imaged with a Zeiss LSM700 confocal scanning microscope (upright), equipped with a 20× water immersion objective (W "Plan-Apochromat" 20×/1.0 DIC M27 75 mm; Zeiss, catalog no. 421452-9800). We consistently used HCR hairpins that were fluorescently labeled with Alexa Fluor 546 and Alexa Fluor 647, and therefore, imaging was performed with the following lasers: Diode 488 (10.0 mW), Diode 555 (10.0 mW), and Diode 639 (5.0 mW). Although we did not experience photobleaching for the majority of samples, some reduction in the intensity of the Alexa Fluor 647 occasionally occurred. To overcome the reduced signal in the ventral parts of the brain due to scattering, we applied z brightness correction, by increasing the laser power in the deep parts of the sample. Z-stacks, comprising two tiles, were scanned in an upward direction, with a dwell time of 1.58 μ s, and stitched to

produce a final image with a size of 1039 \times 1931 pixels (463.97 \times 862.29 $\mu m,$ 1 μm in z).

Generation of reference brain and image registration and averaging

All the image registrations and averaging procedures were performed using ANTs (26), which were previously calibrated and used for the generation of zebrafish average brain images (20, 59). Twelve transgenic fish expressing the transgenes Tg(elavl3:H2b-GCaMP6s) and Tg(elavl3:lynTag-RFP) were fixed and underwent the entire HCR buffer procedure, but without the usage of any probes or hairpins, and imaged as described. The images were used to generate an unbiased average brain by running the following ANTs command on the GWDG computing cluster:

antsMultivariateTemplateConstruction2.sh -d 3 -o \${output-Path}/T_ -i 4 -g 0.2 -j 32 -v 500 -c 2 -k 2 -w 1x1 -f 12x8x4x2 -s 4x3x2x1 -q 200x200x200x200 -n 0 -r 1 -l 1 -z \${target1} -z \${target2} -m CC[2] -t SyN[0.05,6,0.5] \${inputPath}

To register the newly generated reference to the mapzebrain coordinate system, we first registered the older Tg(elavl3:H2b-GCaMP6s) average image that existed in mapzebrain (20) onto the newly generated reference and applied the inverse transformation on the newly generated reference image to align to the mapzebrain coordinate system. First, the following ANTs command was used to transform the older mapzebrain Tg(elavl3:H2b-GCaMP6s)average to a new reference:

antsRegistration -d 3 --float 1 -o [$\{output1\}, \{output2\}\}$ --interpolation WelchWindowedSinc --use-histogram-matching 0 -r [$\{template\}, \{input1\}, 1]$ -t rigid[0.1] -m MI[$\{template\}, \{input1\}, 1]$ -t rigid[0.1] -m MI[$\{template\}, \{input1\}, 1, 32, Regular, 0.25]$ -c [200x200x0, 1e-8, 10] --shrink-factors 12x8x4x2 --smoothing-sigmas 4x3x2x1vox -t Affine[0.1] -m MI[$\{template\}, \{input1\}, 1, 32, Regular, 0.25]$ -c [200x200x200x0, 1e-8, 10] --shrink-factors 12x8x4x2 --smoothing-sigmas 4x3x2x1 -t SyN[0.05, 6, 0.5] -m CC[$\{template\}, \{input1\}, 1, 2]$ -c [200x200x200x200x10, 1e-7, 10] --shrink-factors 12x8x4x2x1 --smoothing-sigmas 4x3x2x1x0

Second, inverse transformation was applied to the newly generated reference using the inverse Warp transformation file by running the following ANTs command:

antsApplyTransforms -d 3 -v 0 -- float -n WelchWindowedSinc -i \${input3} -r \${template} -o \${output4} -t \${output1}1Inverse-Warp.nii.gz -t \${output1}0GenericAffine.mat

The newly registered reference brain (fig. S1 and movie S1) is available to download through https://mapzebrain.org. All the imaged HCR in situ images were registered onto this new reference by registering the Tg(elavl3:H2b-GCaMP6s) image channel using the following ANTs command:

antsRegistration -d 3 --float 1 -o [$\{0,1\}, \{0,1\}, \{0,1\}, \{0,1\}, \{0,1\}, \{1,2\}, \{1,3\},$

followed by applying the transformation files on the HCR image channels using the following ANTs command:

antsApplyTransforms -d 3 -v 0 -- float -n WelchWindowedSinc -i \${input3} -r \${template} -o \${output4} -t \${output1}1Warp.nii.gz -t \${output1}0GenericAffine.mat

Each HCR-labeled RNA was imaged from three different fish and registered as described. The three images were then arithmetically averaged using ImageJ (60). Both the average image and the individual registered images are accessible through https:// mapzebrain.org.

cfos FISH experiments

For *cfos* experiments, larvae were raised until 6 dpf under optimal conditions (14-hour light/10-hour dark cycle, 28°C) in Danieau's solution. At 6 dpf, the larvae were transferred into a six-well plate, 10 larvae per well until 7 dpf, when they were exposed to different stimuli for 40 min. The larvae were then euthanized in an overdose of Tricaine for 2 min and immediately fixed in ice-cold 4% PFA. Six fish of each condition were imaged and registered as described before, followed by arithmetical averaging with ImageJ (*60*). For visualization purposes of the averaged images, the contrast was linearly enhanced. Movie S2 anatomical legend was generated using QuPath (*61*).

Temperature adaptation

The six-well plate containing the fish was placed into a water bath set to either 22° or 32°C with a similar background color for 40 min. *Background adaptation*

The six-well plate containing the fish was placed on either a white or a black background at 28°C for 40 min.

Prey capture

Paramecia were added into the six-well plate containing the larvae. Food was not limited: Many paramecia were still uneaten after 40 min.

Phototaxis

The six-well plate containing the fish was placed on a half-white and half-black background (splitting the well background color equally), at 28°C for 40 min.

Looming

Larvae were exposed to black looming disc stimuli using a setup previously described (62, 63). Briefly, groups of 10 larvae were placed into a 10-cm watch glass containing Danieau's solution and left for 2 hours to acclimate to room temperature and white projector illumination from below. For stimulation, expanding black discs were projected to the center of the dish from below. Discs expanded linearly from 0 to 10 cm diameter within 500 ms, once every 60 s. This cycle was repeated over 40 min.

Optomotor response

Larvae were exposed to moving gratings using a setup previously described (55, 56). Briefly, groups of 10 larvae were placed into a 10-cm watch glass containing Danieau's solution and left for 2 hours to acclimate to a background of a stationary grating (black/ white step grating, 10-mm spatial frequency) projected from below. After 2 hours, gratings were moved in alternating directions at 1 cycle/s in the following sequence: 15 s in one direction, 5 s pause, and 15 s in the opposite direction. This sequence was repeated over 40 min. Visual inspection confirmed a strong behavioral optomotor response under these stimulus conditions. A control group of larvae remained on stationary gratings the entire time.

Alarm substance response

Alarm substance is stored in the club cells of zebrafish epidermis and released upon skin damage. For the experiment, five adult wild-type donor fish were euthanized, and scales from both flanks were collected with the blunt side of a sterile razor blade. The scales were washed off with sterile-filtered Danieau's solution and collected in a Falcon tube, avoiding any contamination with blood. The total 20 ml of isolate was vortexed, filtered through a Whatman paper, and placed on ice until use (64). Five hundred microliters of the isolate was distributed evenly over the wells of the six-well plate containing the larvae.

Single neuron morphology clustering

An ROI surrounding the SGN cfos ISH-labeled area was labeled using the ImageJ segmentation editor plugin and saved as a binary mask .tif file. The .tif file was then uploaded as an ROI to mapzebrain.org and used to search the mapzebrain atlas for neurons whose somas were located within it (to enable this feature in mapzebrain.org, a free user account is required. See movie S3 for a detailed tutorial). Thirty-two neurons were identified and flipped to the left [as described in (20)], and a morphology similarity score between them was calculated using the NBLAST algorithm (51) by applying the nblast_allbyall function implemented in the R package NeuroAnatomy Toolbox (nat) (65) with default parameters. Next, unsupervised hierarchical clustering was performed using the nhclust function of the nat package with default parameters, and the results were plotted using the R packages dendroextras and rgl. The analysis and plotting R code can be found at https:// zenodo.org/record/7308624.

Supplementary Materials

This PDF file includes: Figs. S1 to S3 Table S1 Legends for movies S1 to S3

Other Supplementary Material for this manuscript includes the following: Movies S1 to S3

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