Expansion microscopy of zebrafish for neuroscience and developmental biology studies

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Expansion microscopy (ExM) allows scalable imaging of preserved 3D biological specimens with nanoscale resolution on fast diffraction-limited microscopes. Here, we explore the utility of ExM in the larval and embryonic zebrafish, an important model organism for the study of neuroscience and development. Regarding neuroscience, we found that ExM enabled the tracing of fine processes of radial glia, which are not resolvable with diffraction-limited microscopy. ExM further resolved putative synaptic connections, as well as molecular differences between densely packed synapses. Finally, ExM could resolve subsynaptic protein organization, such as ring-like structures composed of glycine receptors. Regarding development, we used ExM to characterize the shapes of nuclear invaginations and channels, and to visualize cytoskeletal proteins nearby. We detected nuclear invagination channels at late prophase and telophase, potentially suggesting roles for such channels in cell division. Thus, ExM of the larval and embryonic zebrafish may enable fundamental studies of how molecular components are configured in multiple contexts of interest to neuroscience and developmental biology.

Significance

We explore the utility of expansion microscopy (ExM) in neuroscience and developmental biology using the zebrafish model. Regarding neuroscience studies, ExM enables the tracing of cellular processes in the zebrafish brain, as well as the imaging of synapses and their biomolecular content and organization. Regarding development, ExM enables the resolving of nuclear compartments, particularly nuclear invaginations and channels, and helps relate such cellular nanostructures to proteins of the cytoskeleton during embryogenesis.

Methods

Fish Maintenance and Care. All zebrafish (Danio rerio) larvae were raised in fish facility water at Harvard University according to protocols and procedures approved by the Harvard University/Faculty of Arts & Sciences Standing Committee on the Use of Animals in Research and Teaching (Institutional Animal Care and Use Committee), with the following exceptions: Larvae used for imaging were raised in fish facility water at the University of Cambridge. Posthatch larvae were transferred to the facility water at Harvard University according to protocols and procedures approved by the Harvard University/College of Arts & Sciences Standing Committee on the Use of Animals in Research and Teaching.

Conflict of interest statement: E.S.B. and R.G. are inventors on one or more patent applications related to expansion microscopy (ExM). E.S.B. is co-founder of a company, Expansion Technologies, that aims to provide ExM kits and services to the community.

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in Fig. 1 and SI Appendix, Fig. S1 A, B, and D and one of two larvae used for the analysis shown in SI Appendix, Fig. S1C were raised in Danieau’s medium at the Max Planck Institute of Neurobiology. These animal procedures conform to the institutional guidelines of the Max Planck Society and the local government (Regierung von Oberbayern). Experimental protocols were approved by Regierung von Oberbayern (55.2-1-54-2532-101-12 and 55.2-1-54-2532-31-2016). All larvae were raised on a standard 14-h light/10-h dark cycle at a temperature of 28 °C.

Transgenic Fish Lines. The genotypes of the larvae and embryos used to generate each figure are detailed in SI Appendix, SI Methods. The transgenic fish lines that were crossed to produce these larvae and embryos were all previously described. All larval brain images are from 6-d postfertilization larvae, and all embryo images are from shield-stage (i.e., ~6-h postfertilization) embryos.

Immunohistochemistry. Immunohistochemistry was performed following standard, previously published procedures (28). The following reagents were used (29), as well as a detailed list of antibodies used, is provided in SI Appendix, SI Methods.

Expansion. Expansion was performed using the previously described proExM protocol (2) (SI Appendix, SI Methods).

Imaging. Both pre- and postexpansion brains and embryos were imaged on an Andor spinning disk (CSU-X1 Yokogawa) confocal system with a 40×, 1.15 N.A. water immersion objective (Nikon), with the exception of some images in SI Appendix, Fig. S1: The first and third images in SI Appendix, Fig. S1A and all preexpansion images used to generate SI Appendix, Fig. S1C were acquired using a DeltaVision OMX Blaze (GE Healthcare) structured-illumination microscope (SIM) with a 60×, 1.42 N.A. oil immersion objective (Olympus). These brains were immersed in SlowFade Diamond Antifade mounting medium (Invitrogen) for refractive index matching and suppression of bleaching. The first and third images in SI Appendix, Fig. S1B and all preexpansion images used to generate SI Appendix, Fig. S1D were acquired using a Leica TCS SP8 STED microscope, with a 100×, 1.4 N.A. oil immersion objective. These brains were immersed in SlowFade Gold Antifade mounting medium (Invitrogen) for refractive index matching and suppression of bleaching. The second and fourth images in SI Appendix, Fig. S1B and all postexpansion images used to generate SI Appendix, Fig. S1D were acquired using a Leica TCS SP8 confocal microscope, with a 40×, 1.1 N.A. water immersion objective. Images of the samples were also obtained with a 10×, 0.45 N.A. air objective and used to aid in the comparison of pre- and postexpansion data and computation of expansion factors. Details of excitation and emission collection are provided in SI Appendix, SI Methods. For embryos, postexpansion imaging was also performed via a Nikon Ti-E epifluorescence microscope with a 4×, 0.13 N.A. air objective to allow capturing of the entire sample for computation of expansion factors. For brains, expansion factors were computed by measuring the size of specific anatomical features (e.g., the axon cap) pre- vs. postexpansion, and taking the ratio of the respective sizes. For embryos, the diameter of the embryo pre- vs. postexpansion was compared. Scale bars on postexpansion images reflect these expansion factor computations. For the expanded embryos imaged in SI Appendix, Fig. S10, an expansion factor of 4 [similar to the expansion factors computed for other embryos (3.8 and 4.1)] was estimated for the purpose of drawing scale bars.

Image Processing. Each figure panel constitutes a single plane from a z-stack, where the area of interest was cropped out of the field of view using Fiji (29), or a maximal intensity projection, as indicated in the figure legends. The brightness and contrast of individual channels were adjusted in ImageJ (NIH) after cropping the area of interest. The STED preexpansion images shown in SI Appendix, Fig. S1B (first and third panels) and used in SI Appendix, Fig. S1D were deconvolved using Huygens (Scientific Volume Imaging). Tracing of cellular processes (shown in SI Appendix, Fig. S2) was performed using Imaris. This tracing algorithm is intensity-based. First, start and end points of extratectal fibers preexpansion (D) and postexpansion (D′) from the regions highlighted by arrowheads in B and B′, respectively, is shown. (E and E′) Intensity plots along the orange line in D and D′, respectively. AU, arbitrary units. [Scale bars: B, 10 μm; B′, 10 μm (physical size postexpansion, 35 μm); C and D, 5 μm; C′ and D′, 5 μm (17.5 μm).]
Fig. 2. ExM analysis of synaptic connections. (A) Schematic of larval zebrafish brain showing nII and nIV nuclei, labeled by Tg isl1:GFP;Tg(UAS:Kaede) (Kaede, magenta). The rectangular area is imaged in B. (B) Maximal intensity projection of an ∼33-μm-thick volume corresponding to the rectangular area shown in A. The fish is 6 d postfertilization (dpf), and is stained with anti-GFP (yellow), anti-Kaede (magenta), and anti-pan-MAGUK (not shown). (C) GFP-labeled cells (yellow) and Kaede-labeled projections (magenta) in the nII nucleus. (C, I–IV and I–IV′) Two nearby planes (one in each row) from an expanded 5-μm brain stained with anti-GFP (yellow), anti-Kaede (magenta), and anti-synaptotagmin2b (cyan). Arrows point to Kaede-expressing and synaptotagmin2b-stained varicosities and terminals next to GFP-labeled neuropil (IV) and cell bodies (IV′). Arrowheads point to a cluster of synaptotagmin2b, unlabeled by Kaede, next to a GFP-labeled cell body (IV′). (C, V–VIII) Single plane from a brain stained with anti-pan-MAGUK (cyan). Arrows point to Kaede-labeled varicosities and terminals next to GFP-labeled cells and neuropil, exhibiting colocalized MAGUK puncta. Arrowheads point to a MAGUK punctum on a GFP-negative cell opposite to a Kaede-labeled terminal (Top arrowhead) and to MAGUK puncta on GFP-labeled cell bodies and neuropil in the absence of nearby Kaede-labeled projections (Bottom two arrowheads). Scale bars: B, 10 μm (38 μm); C, I–IV and I–IV′, 5 μm (23 μm); C, V–VIII, 5 μm (19 μm).
with the optically clear (1) postexpansion samples, causing reduced signal-to-noise ratios that may obscure some features in preexpansion samples. Finally, orientation differences (even ones remaining after image registration) result in pre- and postexpansion images representing overlapping, yet distinct, optical sections in the samples.

Fig. 3. Expansion enables the resolving of synaptic heterogeneity and structure in intrasynaptic protein distributions. (A) Schematic of a larval zebrafish brain showing the M cells (blue) and spiral fiber neurons (magenta). The rectangle illustrates the region focused in on in B–D, consisting of the axon cap and a part of the M cell body. (B) Preexpansion images of the axon-cap area showing spiral fiber neurons (magenta) wrapping around the M cell axon initial segment (the unlabeled “tube” passing through these fibers, better visualized as a black stripe in C), as well as synaptotagmin2b (Top, cyan) and glycine receptors (Bottom, cyan). (C) Same as in B, but postexpansion. (Note: The synaptotagmin2b axon cap shown (Top) is not from the same brain as in B, Top.) (Top Left and Right) Arrows point to a Kaede-labeled varicosity bearing synaptotagmin2b at a low density. (Top Left and Right) Arrowheads point to a Kaede-negative varicosity bearing dense synaptotagmin2b staining. (Center) Arrowheads point to varicosities in spiral fiber neuron projections forming the M cell axon cap. (D) Maximal intensity projection of the M cell body and axon initial segment area showing the distribution of glycine receptors (cyan) preexpansion (Left) vs. postexpansion (Right). (Note: This is the same axon cap as shown in B and C, Bottom.) Boxes highlight seven examples of ring-shaped clusters zoomed in on in E. (E) Seven examples of ring-shaped clusters of various sizes present on the M cell body (1–6), and axon (7). [Scale bars: B, Top and Bottom, 5 μm; C, Top, 5 μm (23 μm); C, Bottom, 5 μm (20 μm); D, Left, 5 μm; D, Right, 5 μm (20 μm); E, 1 μm (4 μm).]
Nevertheless, a quantitative comparison of the data shows that, in length measurements performed using postexpansion images, the root mean square (rms) of measurement errors relative to the lengths measured was comparable to the results obtained in previously published ExM papers. In particular, the rms of measurement errors was no larger than 5% for preexpansion images acquired with SIM and 2% for preexpansion images acquired with STED microscopy, on average (SI Appendix, Fig. S1 C and D). We note that these estimates constitute higher bounds on actual deformations in the samples, as they are affected by imperfections in the preliminary registration of datasets in three dimensions using rigid transformation and scaling (Methods).

Expansion Facilitates Tracing Cellular Processes in the Zebrafish Tectum. We next set out to assess whether ExM could enable the tracing of fine cellular processes. We used a larva (strain details are provided in Methods) that strongly expresses membrane-bound GFP in a limited number of cells in the tectum (Fig. 1A), and particularly in radial glial cells (Fig. 1B). Expansion made their processes more visible vs. in preexpansion images (compare maximal intensity projections in Fig. 1B′ vs. Fig. 1B and single-plane images of specific regions in Fig. 1C′ vs. Fig. 1C). When we applied commercially available automated tracing software to quantitatively analyze these processes, the number of automatically identified branch points in the traced cells increased approximately sixfold postexpansion vs. when tracing was performed preexpansion (compare SI Appendix, Fig. S2A′ vs. SI Appendix, Fig. S2A and zoomed-in regions in SI Appendix, Fig. S2B′ vs. SI Appendix, Fig. S2B; numbers in the images reflect branch points of the cells indicated; also Movie S1). We were also able to resolve multiple processes within bundles that were not resolvable preexpansion (Fig. 1B′ vs. Fig. 1B, arrowheads and Fig. 1 D′ and E′ vs. Fig. 1 D and E). Thus, ExM may help clarify fine processes in zebrafish cells, as we had earlier found for cellular processes in mouse hippocampus (2).

Resolving Synapses Mediating Sensory/Motor Transformations. We next used ExM to resolve synaptic connections, using larvae (Methods) in which the labeled neurons include some of the key neurons involved in the vestibulo-ocular reflex. In particular, the fluorophore Kaede (39) was expressed in projects into the oculomotor (nIII) and trochlear (nIV) nuclei, including putative tangential neuron projections (12) important for this reflex; furthermore, GFP was expressed in neurons in the nIII and nIV nuclei (highlighted in Fig. 2 A and B and Movie S2), which consist of extracellular motoneurons (40). We found that ExM revealed fine details of Kaede-labeled projections (compare Fig. 2C and SI Appendix, Figs. S3A and S4, examples 1–12, Bottom with SI Appendix, Figs. S3B and S4, examples 1–12, Top), enabling their terminations, which had the appearance of synaptic varicosities, to be visualized. The presynaptic protein synaptotagmin2b (Fig. 2C, I–IV and I–IV′ and SI Appendix, Fig. S3A, I–IV) was localized to such varicosities, and particularly to their boundaries, consistent with these varicosities being presynaptic terminals (arrows in Fig. 2C, IV and IV′ and in SI Appendix, Fig. S3A, IV point to such varicosities abutting right next to GFP-positive cell bodies and neuropil). Similarly, when we immuno-ostained with antibodies against postsynaptic proteins of the MAGUK class, with a pan-MAGUK antibody (41) (Fig. 2C, V–VIII and SI Appendix, Fig. S3A, V–VIII), postexpansion data clearly revealed MAGUK-positive puncta in GFP-labeled cells (arrows in Fig. 2C, VIII and SI Appendix, Fig. S3A, VIII), which were located in apposition to varicosities of Kaede-labeled projections, and thus were putative postsynaptic targets. The ability to characterize the location of synaptic proteins in regard to cell–cell contacts may help pinpoint fine details of synaptic wiring. In particular, examining the Kaede-labeled terminals abutting next to GFP-positive cells in preexpansion data from this pan-MAGUK–stained brain, we identified 12 example terminals where the data suggested more than one GFP-positive cell as a putative postsynaptic target (SI Appendix, Fig. S4, Top). However, examining the same regions in the same brain postexpansion, we found that MAGUK puncta were often present only on a subset of these putative postsynaptic targets, perhaps indicating a selectivity of functional contact not visible in preexpansion data (SI Appendix, Fig. S4, Bottom). The number of GFP-positive targets identified pre- vs. postexpansion for each terminal is indicated in SI Appendix, Table S1. This number is lower in six of the 12 examples in postexpansion compared with preexpansion data. Thus, this analysis suggests that ExM enables more specific assignment of putative postsynaptic partners in the larval zebrafish nervous system.

Expansion Allows Resolving of Subsynaptic Structures. We next explored how ExM could reveal heterogeneity of synaptic composition in dense neuropil, focusing on synapses onto the M cell, which triggers a fast-escape response by firing a single action potential (42, 43). The axon cap is a dense area of neuropil that surrounds the axon initial segment and hillock of the M cell (44) (Fig. 3 A and B, Top Right). To visualize this area, we used larvae (Methods) in which the spiral fiber neurons whose projections form the axon cap (15, 45) express the fluorophore Kaede (46). Putative presynaptic terminals (as indicated with anti-synaptotagmin2b staining) were visible throughout the cap in both preexpansion (Fig. 3B, Top) and postexpansion (Fig. 3C, Top) data, consistent with electron microscopy data indicating the existence of synapses between spiral fiber neurons and themselves, as well as from these neurons onto the M cell axon (15, 47, 48) in the cap. However, individual putative synaptic varicosities were only resolvable postexpansion (Fig. 3C, Center, arrowheads and Movie S3). Furthermore, different synaptotagmin2b-positive varicosities exhibited different amounts of synaptotagmin2b staining (compare arrow vs. arrowhead in Fig. 3C, Top Right and Left), reflecting either synapse-to-synapse variability within a single type of synapse or the presence of multiple molecularly distinct synapse types in the axon cap (15, 47, 48). This heterogeneity also held for the synaptic vesicle glycoprotein 2a (SV2; arrows and top arrowhead vs. bottom arrowhead in SI Appendix, Fig. S4A, Bottom) and the presynaptic proteins synapsin1/2 (SI Appendix, Fig. S4B, Right, arrow vs. arrowhead). To examine whether preExM could go beyond the analysis of synaptic composition and help with the analysis of synaptic structure, we examined the distribution of glycine receptors on the M cell (Fig. 3 B and C, Bottom). These synapses were present on the M cell body and axon, but absent from the rest of the cap. Using a larva that additionally expressed GFP under the control of the glycine transporter-2 (glt2) promoter (49), we found that postexpansion, “log”-like protrusions from glycnergic neurons that approached glycine receptor patches on the M cell were visible (SI Appendix, Fig. S5C, Bottom Right, arrows). Furthermore, glycine receptors formed ring-like patches that were not resolvable preexpansion (Fig. 3D, Right and Movie S4 vs. Fig. 3D, Left). The sizes of these rings varied, with a gradient of decreasing size along the soma toward the axon (Fig. 3E). These findings are consistent with earlier findings in goldfish (16). With ExM, these rings, which are much smaller in the genetically tractable zebrafish, become visible.

Finally, these data also suggested that ExM may enable the better identification of glycine receptor clusters on such putative synapses (SI Appendix, SI Methods). To examine this, we segmented images of four M cell axons in two brains with labeled glycine receptors (SI Appendix, Fig. S6A) and counted the number of distinct receptor clusters as a function of the intensity threshold used in this segmentation. When segmentation was performed with low-intensity thresholds, clusters that visually seemed distinct were merged together, while at high thresholds, some clusters were eliminated. Thus, a peak in the identified number of clusters occurred at an intermediate threshold, representing a trade-off between these two alternatives. Importantly, this analysis revealed that at the peak and...
for a broad range of thresholds around it, the number of distinct clusters identified in postexpansion images was at least sixfold higher than in preexpansion images (SI Appendix, Fig. S6B). In preexpansion images, a low signal-to-noise ratio and limited resolution caused distinct clusters to merge together and small clusters to blend with the background. Thus, this analysis demonstrates that ExM data may be beneficial in counting distinct putative synapses.

**Expansion Reveals the Shape of Intranuclear Invaginations in Embryos.**

To explore ExM in the context of zebrafish developmental biology, we investigated the morphology of nuclei at an early embryonic stage. We used embryos from a fish line where EGF3 is fused to histone-2B (50) (details are provided in Methods), and we used anti-lamin B and anti-tubulin to visualize the nuclear boundary and microtubules, respectively. Of ~350 nuclei across two ~6-h-old zebrafish embryos (~250 and ~100 nuclei in the two embryos, respectively), we found that ~35 nuclei were between the prometaphase and telophase stages, whereas the other ~315 nuclei were in interphase or prophase (i.e., had intact nuclear envelopes). Preexpansion data revealed lamin B within the nucleus, overlapping with labeled histones (Fig. 4A). Postexpansion, such lamin B staining could be resolved as nuclear envelope invaginations containing a cytoplasmic core (i.e., void of histone staining), as had been previously reported in interphase cells (20, 26) (Fig. 4B, Top and Movie S5). Indeed, with ExM, we could see highly variable organizations of such invaginations, ranging from just one or two channels passing through the nucleus (Fig. 4B and SI Appendix, Fig. S7, A and C, I–IV and I) to complex networks of intranuclear channels (SI Appendix, Fig. S7 B and C, V–VII, VI, and VII).

To compare the extent to which the structure of such intranuclear channels could be quantitatively captured using pre- vs. postexpansion images, we sampled 30 nuclei from each dataset (SI Appendix, SI Methods and Figs. S8 and S9). For each nucleus in these samples, the number of disconnected channel structures identified within the nucleus, the number of exit points from the nucleus, and the number of internal end points within the nucleus are enumerated in SI Appendix, Tables S2 and S3 for preand postexpansion samples, respectively. In preexpansion data, it was sometimes difficult to establish whether structures were connected or disconnected, and whether a channel reached all of the way to the nuclear boundary or ended within the nucleus, close to its boundary. Consequently, to capture multiple possible interpretations of the data, we estimated both minimal and maximal values for the above-described features, given the possible image interpretations (SI Appendix, Table S2). We found that, due to the enhanced resolution, complex structures were better captured in postexpansion data, leading to higher estimates for the mean number of disconnected structures and the mean number of exit points identified in nuclei when comparing postexpansion estimates with the minimal preexpansion estimates. Higher exit point counts were found in postexpansion data as well, compared with maximal preexpansion estimates. The number of end points within the nucleus was similar preand postexpansion. Put together, this analysis shows that ExM improved the ability to capture the complexity of intranuclear invagination structures.

We were further able to map out microtubule organization in and around nuclear invaginations, which is of interest since many kinds of cytoskeletal filaments have been associated with such invaginations (20, 22, 24, 26, 27, 51). For example, many centrosomes (clusters of microtubules highlighted with arrowheads in Fig. 4 and SI Appendix, Fig. S7) were localized at invagination openings, although some were offset from the openings (e.g., SI Appendix, Fig. S7, VII, arrow). We also observed microtubule staining at the boundaries of invaginations (Fig. 4B, Bottom and SI Appendix, Fig. S10), although such staining was only visible in a small number of nuclei and the microtubules did not appear continuous. One possibility is that these samples had poor antibody staining, since the dye we used (Methods) was one known to be poorly retained in proExM (2). An additional experiment performed supported this possibility: In two supplementary specimens labeled with dyes known to persist well in proExM (SI Appendix, Fig. S10 and Movie S6), anti-tubulin staining was of higher quality postexpansion. In ~650 nuclei from these two embryos (~400 and ~250 nuclei, respectively), approximately half exhibited microtubules throughout the lengths of nuclear channels (which may still be an underestimate, as the tubulin antibody did not stain samples evenly; Methods). In summary, microtubule staining along the boundaries of intranuclear channels appears to be a widespread phenomenon in shield-stage embryonic nuclei.

**Expansion Reveals Intranuclear Channels at Late Prophase and Telophase.**

Of the ~350 nuclei we analyzed in our main dataset, we found 16 cells (Fig. 5 and SI Appendix, Figs. S7 and S11) in late prophase, with centrosomes on opposing sides of the cell. At this stage, previous studies in cultured mammalian cells suggested that microtubule-containing nuclear envelope indentations form next to centrosomes as the nuclear envelope begins to disassemble (21, 52–56; reviewed in ref. 57.) Indeed, four of the 16 nuclei exhibited indentations ending within the nucleus and starting next to centrosomes (SI Appendix, Fig. S11B, V–VII). However, in 12 of the 16 nuclei, a channel passed through the nucleus, connecting the two centrosomes (one example is shown in Fig. 5A and Movie S7; further examples are shown in SI Appendix, Figs. S7C, VI and VII and S11; arrowheads in all figure panels point to centrosomes). Microtubule staining was sometimes visible within the channel (Fig. 5A, I). Such channels were not resolved preexpansion (although hints were visible; Fig. S5B). We further observed that such nuclear channels might link cytoskeletal
components in unanticipated ways in cells in late telophase. In particular, in late-telophase nuclei (nine of the ∼350 total nuclei; Fig. 5C and SI Appendix, Fig. S12 and Movie S8), we identified intranuclear channels running through daughter nuclei. A centrosome was positioned at one opening of each of these channels (Fig. 5C, I and IV, arrowheads), and microtubules passed through the channels (Fig. 5C, I and IV′), converging at the midbody (Fig. 5C, II, arrow); microtubule staining is absent from the midbody center, likely due to the density of proteins causing epitope masking (58). Thus, these nuclear channels may permit microtubule connections to be made between the now-distant centrosomes. Heterogeneity in the shape and appearance of these structures was apparent from cell to cell; for example, the midbodies of SI Appendix, Fig. S12, I, IV, VI, and VIII seem to exhibit only one microtubule bundle emerging from each side of the midbody rather than two. Overall, however, this configuration (schematized in Fig. 5C, Top) suggests that the nuclear channels might mediate a cross-daughter cell mechanical connection at a time during which the reassembled nuclear envelope would otherwise topologically prohibit such a connection. Another possibility is that the daughter cells have simply advanced beyond telophase, and the phenomena of Fig. 5C are similar to the other phenomena we saw in Fig. 5A (as discussed above); this possibility has been commented on in earlier studies of cells immediately after division (19, 21). Future studies can take advantage of the extended scale imaging capacity of ExM to enable nanoscale imaging of dividing cells, hopefully unraveling structural relations between a variety of proteins and cellular compartments participating in this process (59, 60).

ExM Facilitates Analyses of Mitosis in Zebrafish Embryos. We examined cells in other stages of cell division as well. Post- and preexpansion data from mitotic cells were highly consistent (e.g., compare Fig. 6A vs. Fig. 6B), with some stages nearly identical in appearance (e.g., metaphase images in Fig. 6A, IV vs. Fig. 6B, II′; late anaphase images in Fig. 6A, VII and VIII vs. Fig. 6B, IV and V). However, the structure of microtubules was more accurately captured in postexpansion data vs. preexpansion data, with putative kinetochore-attached microtubule bundles (61) observed to traverse longer distances within nuclei post-expansion [e.g., compare before expansion (Fig. 6B, I and II′) vs. after expansion (Fig. 6A, II′ and III′)], and even the fine and short microtubules of prometaphase were visible postexpansion (Fig. 6A, I′, arrowheads). Furthermore, microtubules near the nucleus boundary (as demarcated with lamin B staining) were easily identified postexpansion in four of five prometaphase nuclei identified in our dataset (Fig. 6A, II′, arrowheads and SI Appendix, Fig. S13, yellow arrowheads; the fifth nucleus was excluded since it was oriented along the optical axis of the objective lens, with its poorer resolution in this direction). Even fine microtubules at early anaphase (Fig. 6A, VI, arrowheads) were observable postexpansion. These various structures are consistent with earlier electron microscopy studies (62, 63), with the exception of the observation of microtubules at nuclear boundaries, which were not described in these earlier studies, perhaps since these studies focused on the internal part of the nucleus and did not explore the boundaries. Similar to the microtubules, lamin B patches were easier to resolve postexpansion (e.g., Fig. 6A, III′, arrowheads) than preexpansion (Fig. 6B, I and II). In particular, patches of lamin B were observable postexpansion to be associated with microtubules that protrude into the nucleus during prometaphase (Fig. 6A, II′′ and SI Appendix, Fig. S13, white arrowheads). Finally, we examined chromatin postexpansion, using the common strategy of imaging EGFP fused to histones as a proxy for chromatin density (64, 65). Postexpansion data recapitulated discoveries made using diffraction-limited light microscopy (66–68). For example, chromatin appeared less densely packed at prophase and early prometaphase (Fig. 6A, I and II) than in late prometaphase and metaphase (Fig. 6A, IV′, V′, and III′), and distinct chromosome arms are not apparent at late anaphase when chromatin is very densely packed (66) (Fig. 6A, VII and VIII). In postexpansion data, chromatin appears (at least in some cell cycle phases) as discrete spots, as in other superresolution microscopy studies (69–71), which may convey its degree of compaction more directly than intensity changes. This suggests that in combination with DNA FISH, it may be possible

Fig. 5. ExM reveals intranuclear channels in late-prophase and telophase nuclei. (A) Nucleus at late prophase, preexpansion. (A, I–III) Three planes from the nucleus. (A, I) Arrowhead points to a centrosome at one opening of an intranuclear channel. (A, II) Arrowhead points to a centrosome at the other opening of the intranuclear channel. (A, IV–VI) Maximal intensity projection of the intranuclear channel-containing area, showing that the channel runs between the two centrosomes located at opposing edges of this nucleus. (B) Nucleus at late prophase, preexpansion, containing two centrosomes at opposite ends of a lamin B-stained channel. (B, I) Representative plane from the nucleus. (B, II–IV) Maximal intensity projection of the invagination-containing region of the nucleus. (C) Structure of a late-telophase nucleus, postexpansion. (Top) Schematic of the structure observed in 10 late-telophase nuclei (eight more nuclei are shown in SI Appendix, Fig. S12). Chromatin (green) is unpacked and fills the daughter nuclei that are surrounded by continuous lamin B staining (blue). Two microtubule bundles (red) diverge from their convergence point at the midbody toward each daughter nucleus. (C, I) Arrow points to a dip in the intensity of microtubule staining at the midbody. (C, IV) Arrowhead points to a centrosome at the channel opening within the second daughter nucleus. [Scale bars: A, 5 μm (19 μm); B, 5 μm; C, 5 μm (20.5 μm).]
to use expansion to map out the location of specific genes in the context of higher level chromatin features.

Discussion

In this study, we primarily compare our data with published electron microscopy studies of larval zebrafish brains and embryos (or cells from other species) rather than with superresolution microscopy imaging studies. This is because, presumably due to the limited z-depth accessible to classical superresolution methods, only a few studies have performed superresolution imaging in zebrafish to date. Techniques used included SIM (72–74), providing a resolution of ∼140 nm and depth penetration of a diversity of rapidly evolving microscopy, providing ∼70-nm resolution but limited depth penetration [14-μm-thick sections of larval zebrafish retina were imaged in ref. 75; a commercial system (Leica TCS SP5 STED) with a reported 90-nm full-width-at-half-maximum lateral resolution and an objective with a working distance of 130 μm was used in ref. 76]. Thus, the successful application of ExM to whole-zebrafish brains and embryos described here may enable the systematic characterization of fine structures in this animal model.

Due to its transparency throughout development, the larval zebrafish is a well-established model for imaging- and optogenetics-based studies of nervous system function (77–79). Synaptic and morphological information provided by ExM would complement such data, and allow linkages to be made between synaptic connectivity and function. In particular, expanding brains after monitoring or manipulating activity in specific cell groups defined by molecular type or connectivity pattern could provide anatomical insights into how information flows or is transformed in the context of behavior. In the future, ExM may be combined with other markers of cellular membranes or cytosol designed to facilitate tracing (e.g., the “spaghetti monster” fluorescent proteins equipped with multiple immunoeptope tags described in ref. 80). Using multicolor labeling (e.g., with Brainbow constructs, as in refs. 81 and 82) could help disambiguate nearby or touching cells that express different combinations of fluorophores or immunoeptopes, reducing the effort associated with neuron tracing (5, 83, 84). Because ExM fills the sample with water, resulting in a transparent, refractive index homogenized specimen (1), such expanded samples are easily scanned with light-sheet microscopes, as we have demonstrated in the context of RNA imaging in expanded mouse brain specimens (85). Embedding preserved biological specimens in polymer hydrogels for imaging purposes has a long history (86), but the use of evenly synthesized swellable polymers to isotropically expand biomolecules away from each other is particularly useful because it can enhance the power of a diversity of rapidly evolving techniques, as described here. In particular, multicolor labeling facilitates antibody staining of multiple proteins at nanoscale resolution with ExM, which is difficult with electron microscopy (87). We regard it as promising that, in parallel to our development of the proExM variant of ExM, two other groups independently developed protocols related to proExM (33, 88), suggesting that proExM is a robust protocol that is easily implemented and deployed.

ExM could be particularly powerful in conjunction with reverse genetic technologies that have been successfully applied to zebrafish (89–92). For example, the structure of radial glial processes and neural projections could be characterized in mutants with compromised development (93–95) to reveal the roles different genes play in wiring the brain. As another example, the scaffolding protein gephyrin localizes glycine (and GABAA) receptors to inhibitory synapses (96). Using ExM in zebrafish, it would be possible to examine the shapes of glycine receptor clusters in mutants where the functions of genes encoding synapse scaffolding components (97, 98) or of other proteins involved in the regulation of such clustering are compromised (99–101).

In the context of development, expanding zebrafish embryos allowed us to identify intranuclear invaginations previously described in other models (20, 26) and to characterize their relationship with microtubules. Previous studies have identified various cytoskeletal filaments in relation to these compartments (20, 22, 24, 26, 27, 51); ExM may enable such analyses to be done rapidly and systematically. Expanding zebrafish embryos could
also help link these structures to specific functions. For example, such invaginations (in cells from other animals) have been proposed to allow regulation of signaling molecule (e.g., Ca\(^{2+}\)) concentration within parts of the nucleus (102, 103), which, in turn, could support spatial regulation of gene expression (23, 104, 105). ExM could allow for the mapping of epigenetic modifications (e.g., caused by mutations in early development) to pass through them and to connect centrosomes at a point in time when the nuclear envelope may otherwise interfere with such direct connections. Of course, from the observation of structure alone, it is impossible to deduce the functional role of these channels, but such hypotheses could be tested in future studies. Studies in zebrafish have revealed a variety of mutants with cell cycle defects [e.g., ones caused by mutations in early-embryonic genes (107)] or deficiencies in chromosome condensation and organization in both interphase and mitosis (108, 109), which could be investigated with ExM. In addition, since incorrect chromosome segregation in mitosis is a potential cause of genomic instability and cancer (110), ExM in zebrafish could be utilized to relate division errors to the development of malignancies (111, 112).

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Movie S1. Three-dimensional rendering of the volume surrounding the radial glia cell RG1 showing this cell and its processes and, subsequently, an overlay of its tracing.

Movie S2. Run through a stack containing the nIII nucleus, with GFP-labeled cell bodies (yellow) and Kaede-labeled projections (magenta) in the expanded 6-d postfertilization larval zebrafish brain stained with anti–pan-MAGUK (cyan) shown in Fig. 2B.
Movie S3. Run through a stack containing the M cell axon-cap area containing spiral fiber neurons (magenta) in the expanded 6-d postfertilization larval zebrafish brain stained with anti-synaptotagmin2b (cyan) shown in Fig. 3C, Top.

Movie S3

Movie S4. Three-dimensional rendering of the M cell body and axon-cap area in the expanded 6-d postfertilization larval zebrafish brain stained with anti-glycine receptor antibody (gray) shown in Fig. 3D, Right.

Movie S4
Movie S5. Run through a stack containing the nucleus shown in Fig. 4B from an expanded shield-stage zebrafish embryo.

Movie S6. Run through the stack containing the nucleus shown in the top row of Fig. S10 from an expanded shield-stage zebrafish embryo.
Movie S7. Run through the stack containing the nucleus shown in Fig. 5A from an expanded shield-stage zebrafish embryo.

Movie S8. Run through the stack containing the nucleus shown in Fig. 5C from an expanded shield-stage zebrafish embryo.

Other Supporting Information Files

SI Appendix (PDF)
SUPPLEMENTARY METHODS

Transgenic fish lines

Fig. 1, SI appendix, Supp. Figs. 1-2 and Supp. Movie 1: *Etk(E1b:Gal4-VP16)s1101t* (1); *Tg(UAS:loxP-tdTomato_CAAAX-loxP-EGFP_CAAAX)mpn128* (2) embryos, expressing membrane-bound tdTomato throughout the brain, were injected at the 1-cell-stage with 100 ng/µl of a Tol2-based Cre construct that drives expression in most tectal cells, *otx1b:Cre*, together with 25 ng/µl Tol2 RNA to mediate recombination. Cells expressing Cre following the injection are thus labeled with membrane-bound EGFP instead of tdTomato. The *otx1b:Cre* construct was generated by inserting a DNA fragment containing a Cre sequence downstream of the *otx1b* promoter in a bacterial artificial chromosome (BAC) (clone #DKEY-209N21), using standard recombineering techniques (3).

SI appendix, Supp. Fig. 1C: One of the two larvae used to generate this panel was a wild-type larva (the second larva was as described above).

Fig. 2B-C, SI appendix, Supp. Figs. 3-4 and Supp. Movie 2: *Tg(-6.7FRhcrtr:Gal4VP16)* (4); *Tg(UAS:Kaede)* (5); *Tg(isl1:GFP)rw0* (6) larvae, expressing GFP in cranial sensory and motor neurons, and Kaede in spiral fiber neurons as well as in neurons from the tangential and medial vestibular nuclei. Kaede was photoconverted from green to red using UV exposure via an epifluorescent microscope (Olympus MVX10) for a duration of 45 seconds.

Fig. 3B-E and Supp. Movies 3-4: The brain labeled with anti-synaptotagmin2b is from the same larva used for Fig. 2. The brain labeled with anti-glycine receptors is from a *Tg(-6.7FRhcrtr:Gal4VP16); Tg(UAS:GFP)* larva, expressing GFP in the same subset of neurons as described above.

SI appendix, Supp. Fig. 5A: A *Tg(-6.7FRhcrtr:Gal4VP16); Tg(UAS:GFP)* larva.

SI appendix, Supp. Fig. 5B: A wild-type larva.

SI appendix, Supp. Fig. 5C: A *Tg(glyt2:GFP)* (7); *Tg(-6.7FRhcrtr:Gal4VP16); Tg(UAS:Kaede)* larva, expressing Kaede in the same subset of neurons as described above as well as GFP in glycineric (inhibitory) neurons. Kaede was photoconverted from green to red as described above.

SI appendix, Supp. Fig. 6: A, top three rows, data from the same larva as in Fig. 3B-E. A, bottom three rows, data from a distinct larva with the same genotype. B, data from both larvae.

Figs. 4-6, SI appendix, Supp. Figs. 7-13 and Supp. Movies 5-8: *Tg(actb2:h2b-egfp/actb2:mem-mCherry2) (hm31)* embryos (8), over-expressing EGFP bound to histones and membrane-bound mCherry in all cells. Note that while these embryos express both EGFP in the nucleus and
mCherry on the membrane of all cells, we only amplified the former with antibodies (see below), and the expression of the latter was rather weak. Membrane labeling was visible in images of embryos both pre- and post- expansion, yet dimmer than other stains. This labeling of membranes did not interfere with the imaging of spectrally overlapping staining which was localized to the nucleus and cytosol.

**Immunohistochemistry protocol and antibodies used**

Briefly, embryos were dechorionated prior to fixation via treatment with 1 mg/ml pronase (Sigma, P5147) for 5 min at room temperature. 6 days post-fertilization (dpf) larvae were fixed in 4% paraformaldehyde (Electron Microscopy Sciences, PA) in phosphate buffered saline (PBS) for 8 hours at 4°C. Embryos were fixed in 4% formaldehyde (diluted from a 37% stock, BDH, 0500-1LP), overnight at 4°C. After fixation, whole larval brains were manually dissected out of the larvae, removing all the skin around the brains. Prior to primary antibody application, embryos were placed in blocking buffer (2% Normal Goat Serum, 1% Bovine Serum Albumin, 1% DMSO, 0.25% Triton-X in PBS) for at least 2 hours at room temperature, and larval brains for at least 1 hour at 4°C. Antigen retrieval was applied to larval brains by placing them for 5 minutes at room temperature, then for 15 minutes at 70°C, in 150 mM Tris-HCl, pH 9.0 (9). Primary antibodies were diluted 1:100 in blocking buffer and applied for no less than 1 day and up to 3 days at 4°C. Secondary antibodies were diluted 1:200 and applied for the same duration, also at 4°C.

The following primary and secondary antibodies were used to generate figures:

**Fig. 1, SI appendix, Supp. Figs. 1-2 and Supp. Movie 1**: Primary – chicken anti-GFP (Invitrogen, A10262). Secondary – Alexa Fluor 488 goat anti-chicken (Invitrogen, A11039).

**SI appendix, Supp. Fig. 1C**, one of the two larvae used to generate this panel was stained with: primary – mouse anti-α-tubulin (Sigma-Aldrich, T6074); secondary – Alexa Fluor 546 goat anti-mouse (Invitrogen, A11030). (The second larva was stained as described above.)

**Fig. 2B-C, SI appendix, Supp. Figs. 3-4 and Supp. Movie 2**: Primaries – chicken anti-GFP; rabbit anti-Kaede (MBL, PM012M); mouse anti znp-1 (DSHB, znp-1) or mouse anti pan-MAGUK (Antibodies Inc., 73-029, NeuroMab clone K28/82; binds PSD-95, PSD-93 and SAP-97 (10)). Secondaries – In anti-synaptotagmin2b (anti znp-1) stained brains: Alexa Fluor 488 goat anti-chicken; Alexa Fluor 568 goat anti-rabbit (Invitrogen, A11011); CF633 donkey anti-mouse (Biotium, 20124). In anti-MAGUK stained brains: Alexa Fluor 488 goat anti-chicken; Alexa Fluor 546 goat anti-rabbit (Invitrogen, A11035); Atto647N goat anti-mouse (Sigma-Aldrich, 50185).

**Fig. 3B-E, SI appendix, Supp. Fig. 6 and Supp. Movies 3-4**: Same as in Fig. 2 for the brain immunostained with anti-synaptotagmin2b. For the brain labeled with anti-glycine receptors:
Primaries – chicken anti-GFP; mouse anti-glycine receptors (Synaptic systems, 146 011). Secondaries – Alexa Fluor 488 goat anti-chicken; Atto647N goat anti-mouse.

**SI appendix, Supp. Fig. 5A**: Primaries – chicken anti-GFP; mouse IgG2a anti zn-1; mouse IgG1 anti-SV2 (DSHB, SV2). Secondaries – Alexa Fluor 488 goat anti-chicken; Alexa Fluor 568 anti-mouse IgG2a (Invitrogen, A-21134); CF405S goat anti-mouse IgG1 (Biotium, 20380).

**SI appendix, Supp. Fig. 5B**: Primary – rabbit anti-synapsin1/2 (Synaptic systems, 106-002); Secondary – Alexa Fluor 488 goat anti-rabbit (Invitrogen, A11008).

**SI appendix, Supp. Fig. 5C**: Primaries – chicken anti-GFP; rabbit anti-Kaede; mouse anti glycine receptors. Secondaries – Alexa Fluor 488 goat anti-chicken; Alexa Fluor 546 goat anti-rabbit; Cy5 goat anti-mouse (Invitrogen, A10524).

**Figs. 4-6 and SI appendix, Supp. Figs. 7-9, 11-13 and Supp. Movies 5 and 7-8**: Primaries – chicken anti-GFP; rabbit anti-lamin B1 (Abcam, ab16048); mouse anti-α-tubulin. Secondaries – Alexa Fluor 488 goat anti-chicken; Alexa Fluor 546 goat anti-rabbit; Cy5 goat anti-mouse.

**SI appendix, Supp. Fig. 10 and Supp. Movie 6**: Primaries - chicken anti-GFP; mouse anti-α-tubulin. Secondaries - Alexa Fluor 488 goat anti-chicken; Alexa Fluor 546 goat anti-mouse.

**Notes on expansion factors and staining quality pre- and post- expansion**

The larval zebrafish brain shown in Fig. 2CI-IV and I'-IV', Fig. 3B-C, top rows, and Supp. Fig. 3AI-IV was gelled with highly pure sodium acrylate (as judged by its appearance), and a particularly high expansion factor (4.6) was obtained. The larval zebrafish brain shown in Supp. Fig. 5A was accidentally placed in a Ca\(^{2+}\)-containing solution prior to expansion, and consequently did not fully expand (this brain’s expansion factor was 2.98). Data presented in Fig. 3B-E, Supp. Fig. 5A-B and Supp. Fig. 6A reflects very high-quality staining (as judged by post-expansion imaging signal-to-noise ratios), which was obtained in a small fraction of staining attempts performed. We found that particular care must be taken regarding staining conditions when staining synaptic proteins, including the need for long incubation times, and it can be helpful to evaluate staining with individual antibodies before combining them into mixtures; one possibility (although speculative) is that some antibody combinations may have reduced staining density vs. when the antibodies are delivered separately (but the pattern and structure of staining are conserved).

Embryos shown in Figs. 4-6 and Supp. Figs. 7-9 and 11-13 as well as the brain shown in Supp. Fig. 5C were stained with a Cy5-conjugated secondary antibody. This dye has relatively poor retention in proExM (11) and consequently, the corresponding signal in expanded images is weaker than it could have been if a different infrared dye (such as Atto647N or CF633) had been
used. Furthermore, the penetration of mouse anti-α-tubulin (Sigma-Aldrich, T6074) primary antibody into embryos was poor, and therefore we used a high concentration of this antibody. However, shallow cells in embryos were very strongly stained, including some non-specific staining, and microtubules in deep cells were still only weakly stained. In cells at intermediate depth, however, staining was strong and specific. We judged staining quality by looking at dividing cells, where the structure of microtubule fibers is stereotyped; if such dividing cells appeared high-quality, we would select nearby cells (e.g., non-dividing ones) for analysis.

Expansion
Brains already stained with primary and secondary antibodies were treated with 0.1 mg/ml Acryloyl-X, SE (6-((acryloyl)amino)hexanoic acid, succinimidyl ester (Molecular probes, A20770; abbreviated AcX) in PBS, made from a 10 mg/ml stock in DMSO, overnight at room temperature. For gelation, 0.2% w/w ammonium persulfate (APS) initiator and tetramethylethlenediamine (TEMED) accelerator were added to the monomer solution (1x PBS, 2 M NaCl, 8.625% (w/w) sodium acrylate, 2.5% (w/w) acrylamide, 0.15% (w/w) N,N'-methylenebisacrylamide) at a concentration of 0.2% (w/w) each from 10% (w/w) stocks. The inhibitor 4-hydroxy-2,2,6,6-tetramethylpiperidin-1-oxyl (4-HT) was added at a concentration of up to 0.01% (w/w) from a 0.5% (w/w) stock. Gelation was performed in a gel chamber, made similarly to the tissue-slice gel chamber described previously (12), using two #2 coverslips as spacers. This chamber was thick enough to fit both larval brains as well as shield-stage embryos separated from their yolk sacs. Brains and embryos were incubated in the chamber for 1 hour at 4°C to allow monomers to diffuse through the tissues, then transferred to a humidified 37°C incubator for 2 hours for gelation. Digestion was performed by incubating gels overnight at 50°C in Proteinase K (New England Biolabs), diluted 1:50 to 16 units/mL in digestion buffer (50 mM Tris pH 8, 1 mM EDTA, 0.5% Triton X-100, 0.5 M NaCl). Then, gels were expanded by placing them in excess volumes of doubly deionized water for 30 minutes, replacing the water for additional 30-minute washes 4 times.

Imaging: excitation and emission collection wavelengths
For images obtained on the Andor spinning disk confocal system, 3 excitation wavelengths were used: 488 nm, 561 nm and 642 nm. Green fluorescence was collected through a 525/50 nm filter, red through a 617/73 nm filter and far red through a 685/40 nm filter. For images obtained on the SIM, 488 nm and 568 nm excitation wavelengths were used. Green fluorescence was collected through a 528/48 nm filter, and red through a 609/37 nm filter. For images obtained on the Leica TCS SP8 STED microscope, an excitation wavelength of 488 nm and a depletion wavelength of
592 nm were used, and emission collection was between 502 and 561 nm. For images obtained on the Leica TCS SP8 confocal microscope, an excitation wavelength of 488 nm was used and emission collection was between 500 and 550 nm.

**Measurement error quantification**

First, unlike thin mouse brain slices, zebrafish brains, due to their ellipsoid-like morphology, are prone to high variability in their 3D orientation when mounted for imaging and gelling. Consequently, the 3D orientation of data acquired pre- and post-expansion varied significantly, potentially contributing to variability in comparing pre- vs. post-expansion data, and a 3D similarity transform was applied to register the datasets. In particular, we first scaled images by measuring the length of specific features pre- and post-expansion to estimate the expansion factor. Then, Fiji's "Name landmark and register" plugin was used to identify corresponding points in the scaled pre- and post-expansion datasets and apply a rigid transformation to register the two datasets. To generate **Supp. Fig. 1A-B**, we performed each registration twice, first transforming pre-expansion data to match the post-expansion data orientation, and then vice versa. Maximal intensity projections of both transformed and un-transformed pre- and post-expansion image stacks are thus shown. For measurement quantification purposes, we used post-expansion data transformed to pre-expansion data in both orientation and scale. Measurement errors were computed on selected single images out of aligned stacks which contained a high number of features that could be used for registration. To compute a vector deformation field expressing the shift of each point in post-expansion images relative to ideal, isotropic expansion we manually selected correspondence points and used the B-spline based non-rigid registration package in Matlab as before. Relative localization errors in using post-expansion images to measure distances between pairs of points were computed also as before, by sampling the entire population of possible point-to-point measurements, and subtracting vectors from the vector deformation fields at the sampled point pairs. In this manner, the root mean square (rms) error for such measurements was computed as a function of the measured length, as presented in **Supp. Fig. 1C-D**.

**Glycine receptor cluster segmentation**

Glycine receptor clusters on M cell axon images were segmented, as shown in **SI appendix, Supp. Fig. 6**, using code written in Matlab. First, the axon was manually segmented out of each image. Then, intensity in the masked image was normalized to a 0-1 range. An intensity threshold was applied to the resulting image to generate a binary image where all pixels with intensities higher than the threshold were replaced with binary 1’s and the rest with 0’s. The number of
distinct connected components (with a connectivity of 8) in the binary thresholded image was then found. Components that were too small (occupied less than 3 pixels pre-expansion and 12 pixels post-expansion) were not counted, as visual inspection revealed that such small components typically represented noise.

**Analysis of embryonic cell nuclei morphology**
The morphology of embryonic cell nuclei was visually assessed by examining the structure observed in the lamin B staining channel. This was performed on samples of 30 nuclei from neighboring cells selected at random from each of the pre- and post- expansion datasets. For each nucleus, the number of disconnected structures present, the number of exit points from the nucleus, and the number of internal end points within the nucleus, were counted. An unpaired t-test was used to establish statistically significant differences in the mean values of these features.

**Supplemental movies**
The 3D volume rendering for **Supp. Movie 1** was generated using Imaris. **Supp. Movie 2** shows the entire stitched stack (from a brain stained with anti pan-MAGUK, also shown in Fig. 2 and SI appendix, Supp. Figs. 3-4) consisting of the region of interest where putative synapses between tangential neuron projections (magenta) and oculomotor (nIII) as well as trochlear (nIV) nuclei cell bodies (yellow) reside. **Supp. Movie 3** shows the cropped-out stack of the axon-cap region in the expanded brain stained with anti-synaptotagmin2b, also shown in Fig. 3C. 3D volume rendering for **Supp. Movie 4** was generated using Vaa3D (13), after cropping the axon-cap region of the anti-glycine receptor stained brain and masking the data to only show the M cell body and axon. The M cell mask was generated manually, using the “segmentation editor” plugin included in Fiji. **Supp. Movies 5-8** show individual cell nuclei cropped out from stacks containing images of many cell nuclei in expanded embryos.
**Supplementary Tables**

**Supp. Table 1.** Number of post-synaptic GFP-expressing targets identified for individual kaede-expressing terminals in pre- vs. post- expansion data (see Supp. Fig. 4).

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**Supp. Table 2.** Morphologies of intra-nuclear invagination structures in shield-stage embryos annotated using pre-expansion data (see Supp. Fig. 8).

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**Supp. Table 3.** Morphologies of intra-nuclear invagination structures in shield-stage embryos annotated using post-expansion data (see Supp. Fig. 9).

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**SUPPLEMENTARY FIGURES**

**A**

Pre expansion (SIM) transformed

Post expansion

Pre expansion (SIM)

Post expansion transformed

**B**

Pre expansion (STED) transformed

Post expansion

Pre expansion (STED)

Post expansion transformed

**C**

SIM

D

STED

Supp. Fig. 1. Validation of expansion microscopy of zebrafish. 

**A,B**, Pre- and post- expansion images of 6 dpf larval zebrafish brains sparsely expressing membrane-bound EGFP (EGFP-CAAX) in a small subset of cells (see Fig. 1 and related text). Left two images of each row: maximal intensity projections of pre- and post- expansion stacks where a similarity transformation was applied to align the pre-expansion stack with the post-expansion stack. Right two images of each row: the similarity transformation was applied to the post-expansion stack, to align it to the pre-expansion stack. All image panels are maximum intensity projections. 

**A**, Pre-expansion data was acquired with a structured-illumination microscope (SIM), and post-expansion data with a spinning disk confocal microscope. 

**B**, Pre-expansion data was acquired with a stimulated emission depletion (STED) microscope, and post-expansion data with a confocal microscope. 

**C,D**, root-mean square (rms) of length measurement errors of ExM versus SIM images (C, n = 4 regions from 2 different brains) and of ExM versus STED images (D, n = 2 regions from 2 different brains) as a function of the measured lengths. Scale bars: 

**A,B**, 3rd image from the left, 5 µm; 

**A**, 2nd image from the left, 5 µm (physical size post-expansion 16.5 µm); 

**B**, 2nd image from the left, 5 µm (18 µm).
Supp. Figure 2. ExM gives rise to more detailed tracings of fine cellular processes. A,A', The radial glial cells RG1 and RG2 were traced from their cell bodies to their endfeet (green and yellow respectively). The number of identified branch points is as indicated. B,B', Close-up on the end-feet of RG1, showing raw imaging data (left) and tracing (green) overlay (right). Scale bars: A-B, 5 µm; A'-B', 5 µm (17.5 µm).
Supp. Figure 3. ExM analysis of synaptic connections: further examples. **A,** As in Fig. 2C, but for the trochlear nucleus (nIV). **AIV,** Arrows, Kaede-labeled synaptotagmin2b-stained projections next to GFP-labeled cells. Arrowhead, a Kaede-negative synaptotagmin2b cluster next to a GFP-labeled cell. **AVIII,** Arrows, Kaede-labeled projections opposed to a GFP-labeled cell body (top) and neuropil (bottom) co-localized with MAGUK puncta. Arrowhead, a MAGUK punctum co-localized with a GFP-labeled cell body in the absence of nearby Kaede-labeled projections. **B,** Pre-expansion data from the same region of the same brain of **AV-VIII.** Scale bars: **AIV,** 5 µm (23 µm); **AVIII,** 5 µm (19 µm); **BIV,** 5 µm.
Supp. Figure 4. ExM enables more specific identification of putative synaptic partners. Examples of terminals abutting next to more than one cell from the same brain as shown in Fig. 2B, 2CV-VIII, Supp. Fig. 3AV-VIII and 3B. Color code is as in Supp. Fig. 3B. In each example: top row, pre-expansion; bottom row, post-expansion. Arrows, MAGUK puncta overlapping with GFP-labeled cells or neuropil right next to the kaede-labeled terminal, indicating a putative synapse made with a specific post-synaptic partner. In pre-expansion data, planes presented show the variety of distinct cells which the terminal is observed to approach. In post-expansion data, planes presented show putative synapses made, with specific putative post-synaptic partners identified. Scale bars: top rows, 5 µm; bottom rows, 5 µm (19 µm).
Supp. Figure 5. Heterogeneity in synaptic protein localization to synapses within the axon cap area of the M cell.  

A, Pre- (top two rows) and post- (bottom two rows) expansion data from a Tg(-6.7FRhcr::R:Gal4VP16); Tg(UAS:Kaede) 6 dpf larval zebrafish brain stained with both anti-SV2 (top left of each dataset, yellow) and anti-synaptotagmin2b (bottom left of each dataset, cyan). In post-expansion images: Top arrowhead, a Kaede-negative varicosity strongly stained with both anti-SV2 and anti-synaptotagmin2b. Bottom arrowhead, a Kaede-negative varicosity showing weak anti-SV2 and anti-synaptotagmin2b staining. Arrows, Kaede-labeled varicosities strongly stained with both anti-SV2 and anti-synaptotagmin2b. 

B, Pre- (left) and post- (right) expansion images of a 6 dpf wild-type larval zebrafish brain stained with an anti-synapsin1/2 antibody. Arrow, a varicosity densely stained with anti-synapsin1/2. Arrowhead, a varicosity weakly stained with anti-synapsin1/2. 

C, Pre- (top) and post- (bottom) expansion images of the M cell body, surrounded by glycine receptors (cyan), receiving inputs from Kaede (magenta)-labeled neurons and glycinergic neurons (yellow). Arrows: "leg"-like protrusions of glycinergic neurons next to glycine receptor clusters on the M cell body. 

Scale bars: A, pre-expansion panel, 5 µm; post-expansion panel, 5 µm (14.9 µm); B, left, 5 µm; right, 5 µm (21.5 µm); C, top, 5 µm; bottom, 5 µm (20.5 µm).
Supp. Figure 6. ExM enables identifying more distinct putative synapses. A, Pre- (left) and post- (right) expansion images of an M cell axon. Top 3 rows: brain 1 (also shown in Fig. 3B-E); bottom 3 rows: brain 2. In each group of 3 rows: top, maximal projection of the axon. Middle, segmentation result using the minimal intensity threshold that brings the number of distinct identified glycine receptor clusters (putative synapses) to a maximum (see B). Distinct clusters are labeled in slightly different colors. Bottom, the axon image thresholded using the same threshold as was used to generate the segmentation image above. B, The number of clusters (putative synapses) identified as a function of the threshold value used in pre- (blue) or post- (red) expansion data. Scale bars: A, left, brains 1 and 2, 5 µm. Right, brain 1, 5 µm (20.5 µm), brain 2, 5 µm (19.5).
Supp. Figure 7. A variety of intra-nuclear invagination structures observed in zebrafish embryonic cells. In all figure panels in this figure and in Supp. Figs. 8-13: blue, anti-lamin B; green, histone 2B (EGFP fused to histone 2B in Tg(actb2: h2b-egfp/actb2: mem-mCherry2), and then stained with anti-GFP); red, anti-α-tubulin (microtubules). A, A nucleus, post-expansion. Al-III, Three planes from the nucleus. Al, Arrowhead, a centrosome at the convergence point of two channels traversing this nucleus. AIV, Maximal intensity projection of the invagination-containing area of this nucleus. B, A second nucleus, post-expansion, showing a complex channel network consisting of at least five exit points, with centrosomes observed at two. BI-III, Three planes from the nucleus. Arrowheads, centrosomes. BIV, Maximal intensity projection of the invagination-containing area of the nucleus. CI',II-IV', VII', Maximal intensity projections of the invagination-containing area of seven different nuclei from two different shield-stage embryos, post-expansion. CI,VI, VII', Individual planes from the same nuclei. CI, A nucleus containing a single channel passing through it. Arrowhead, a centrosome at the entry point of a channel running through the nucleus. CI, A nucleus containing a single channel that splits into two branches before exiting the nucleus. CI, A nucleus containing a single channel passing through it and a nearby deep indentation that ends within it. CIV, A nucleus containing two disconnected channels passing through it. CV-IV, Nuclei containing complex channel networks. CVI, Arrowhead, a centrosome at one of the entry points of a complex channel network. CVII, Top arrowhead, a centrosome close to the entry point of a complex channel network. Arrow, the external edge of a branch in the complex channel network. Bottom arrowhead, a centrosome positioned away from an entry point of the complex channel network. Scale bars: A,B,CI,CIV-VII, 5 µm (20.5 µm); CI-III, 5 µm (19 µm).
Supp. Figure 8. A sample of intra-nuclear invagination structures in pre-expansion data. Each figure panel is a maximal intensity projection of the invagination-containing area of a nucleus in a shield-stage embryo imaged pre-expansion. Scale bars: 5 µm.
Supp. Figure 9. A sample of intra-nuclear invagination structures in post-expansion data. Each figure panel is a maximal intensity projection of the invagination-containing area of a nucleus in a shield-stage embryo imaged post-expansion. Scale bars: 5 µm (20.5 µm). Samples are not the same as used to generate Supp. Fig. 8.
Supp. Figure 10. Microtubules along the boundaries of intra-nuclear invagination channels. Each row is from a plane of a different cell nucleus from one of two expanded embryos, showing clear microtubule staining (red) at the boundary of an intra-nuclear invagination channel, identifiable by the absence of histone labeling (green) within the channel. Scale bars: 5 µm, assuming an expansion factor of 4 (20 µm)
Supp. Figure 11. Intra-nuclear invagination structures in late prophase nuclei. A-B, Examples of nuclear structures similar to the one shown in Fig. 5A, observed in prophase nuclei of expanded embryos. A, Left column, single plane from the nucleus. Right three columns, Maximal intensity projection of the centrosome and nearby invagination containing region of the nucleus. AI-II, Nuclei containing a channel running between the two centrosomes, which, in contrast to the nucleus shown in Fig. 5A, have no apparent microtubule staining within the channels. Note, however, that this absence of observed microtubule staining may be due to poor staining quality as described in the Results and Methods. AI, Arrowheads, centrosomes at
the two openings of a channel. AII, Right arrowhead, centrosome at one side of a channel opening. Left arrowhead, a centrosome at the edge of a channel opening. AIII, In this nucleus both centrosomes are located in a plane above the one through which a microtubule-stained channel traverses the nucleus. Arrowheads, centrosomes. AIV, In this nucleus, one centrosome is located away from an invagination channel opening while the other is located at an opening to an indentation that ends within the nucleus (the maximal intensity projection reflects the presence of a channel running through the nucleus at a distinct plane). Arrowhead, the centrosome located at an opening of a nuclear-membrane indentation that ends within the nucleus. AV, In this nucleus, one centrosome is located at a channel opening while the other is located at the opening of a nuclear-membrane indentation that ends within the nucleus (the maximal intensity projection reflects the presence of a channel running through the nucleus at a distinct plane). Arrowhead, the centrosome located at the opening of a nuclear-membrane indentation that ends within the nucleus. B, Maximal intensity projections of the areas containing centrosomes and nearby invaginations in 8 more nuclei at late prophase. BI, A nucleus containing a channel running through it in between two centrosomes, located at the edge of the nucleus rather than deep within it as the channel shown in Fig. 5A. BII, A nucleus containing a channel passing through it with one centrosome located at a channel opening and another located near the other channel opening. BIII, A nucleus containing a curved, rather than straight as in Fig. 5A, channel connecting the two opposing centrosomes. BIV, A nucleus containing a branched channel running between the two opposing centrosomes. BV-BVIII, Nuclei containing centrosomes at the openings of indentations ending within the nucleus. Scale bars: Al-II, AlV-BI-VI, BVIII, 5 µm (20.5 µm); AlIII, BVII, 5 µm (19 µm).
Supp. Figure 12. Intra-nuclear channels in telophase cells. In each row, planes of a different late telophase nucleus are presented. V, Same nucleus as shown in Fig. 6AIX. Scale bars: I-VI, 5 µm (20.5 µm); VII-VIII, 5 µm (19 µm).
Supp. Figure 13. Lamin B staining near microtubules in prometaphase nuclei. Microtubules (red) and lamin B (blue) staining in 3 selected planes within 4 different nuclei (one in each row) at prometaphase. White arrowheads, lamin B staining next to microtubules that protrude into the nucleus. Yellow arrowheads, microtubules at the boundaries of nuclei, overlapping with surrounding lamin B staining. I-III. Prometaphase nuclei where lamin B staining is observed next to nearly all microtubules that protrude into the nucleus. IV. A prometaphase nucleus approaching the transition from prometaphase to metaphase. Consequently, the number of microtubules protruding into this nucleus was larger than in the other nuclei, lamin B was more depolymerized and chromosomes were presumably more tightly organized near the spindle equator. The fraction of microtubules protruding into the nucleus that were associated with nearby lamin B staining in this nucleus was lower than in the other prometaphase nuclei described above. Scale bars: 5 µm (20.5 µm)
REFERENCES


