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# Citation

Engerer, Peter, Sachihiro C Suzuki, Takeshi Yoshimatsu, Prisca Chapouton, Nancy Obeng, Benjamin Odermatt, Philip R Williams, Thomas Misgeld, and Leanne Godinho. 2017. "Uncoupling of neurogenesis and differentiation during retinal development." The EMBO Journal 36 (9): 1134-1146. doi:10.15252/embj.201694230. http://dx.doi.org/10.15252/embj.201694230.

# **Published Version**

doi:10.15252/embj.201694230

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# Article



# Uncoupling of neurogenesis and differentiation during retinal development

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### Abstract

Conventionally, neuronal development is regarded to follow a stereotypic sequence of neurogenesis, migration, and differentiation. We demonstrate that this notion is not a general principle of neuronal development by documenting the timing of mitosis in relation to multiple differentiation events for bipolar cells (BCs) in the zebrafish retina using in vivo imaging. We found that BC progenitors undergo terminal neurogenic divisions while in markedly disparate stages of neuronal differentiation. Remarkably, the differentiation state of individual BC progenitors at mitosis is not arbitrary but matches the differentiation state of post-mitotic BCs in their surround. By experimentally shifting the relative timing of progenitor division and differentiation, we provide evidence that neurogenesis and differentiation can occur independently of each other. We propose that the uncoupling of neurogenesis and differentiation could provide neurogenic programs with flexibility, while allowing for synchronous neuronal development within a continuously expanding cell pool.

Keywords bipolar cells; development; differentiation; neurogenesis; retina
Subject Categories Development & Differentiation; Neuroscience
DOI 10.15252/embj.201694230 | Received 1 March 2016 | Revised 17 January
2017 | Accepted 24 January 2017 | Published online 3 March 2017
The EMBO Journal (2017) 36: 1134–1146

## Introduction

The nervous system is indisputably the most complex structure assembled during vertebrate ontogenesis. Consequently, the developmental processes underlying nervous system assembly must be precisely orchestrated. Neuronal development in the central and peripheral nervous systems (CNS, PNS) is widely accepted to require three major steps: (i) neurogenesis, the birth of neurons by progenitor cell mitosis, (ii) migration, the relocation of post-mitotic neurons from their birthplace in proliferative zones to specific locations, and (iii) neuronal differentiation, the acquisition of molecular and morphological features that permit the integration of newly generated neurons into synaptic circuits. To date, the prevailing view is that these ontogenetic events are discrete steps along a stereotypic sequence, beginning with neurogenesis, followed by migration and concluding with neuronal differentiation. There is however, particularly in the PNS, evidence that some hallmarks of neuronal differentiation might already occur in progenitors (Rothman et al, 1980; Rohrer & Thoenen, 1987; DiCicco-Bloom et al, 1990; Godinho et al, 2007; Attardo et al, 2008). These observations call into question the invariant developmental sequence of neurogenesis, migration, and differentiation, and raise the unresolved question: How stereotypic is the developmental program of a defined progenitor population in vivo?

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Here, we examined the developmental fate of a molecularly defined CNS progenitor population (expressing *visual homeobox gene 1, vsx1*) that gives rise to the vast majority of a specific interneuron cell type (bipolar cells, BCs) in the zebrafish retina by terminal symmetric divisions (He *et al*, 2012; Weber *et al*, 2014). The swift development as well as the genetic and optical accessibility of zebrafish permitted us to follow the entire developmental program of  $vsx1^+$  progenitors, with single cell precision, *in vivo*. We used molecular, morphological, and cell biological markers of neuronal differentiation, in conjunction with chronic *in vivo* time-lapse imaging, to determine the timing of mitosis in relation to a battery of developmental events. We discovered that for BCs, neurogenesis and multiple hallmarks of neuronal differentiation (such as

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<sup>[</sup>The copyright line of this article was changed on 2 May 2017 after original online publication.]

somal positioning, neuronal marker expression, or neurite elaboration) are timed independently of each other. In other words, rather than dividing at a stereotypic point in their developmental trajectory,  $vsx1^+$  progenitors of BCs undergo terminal mitosis at markedly disparate stages of differentiation, suggesting that differentiation is not time-locked to mitosis. However, the state of differentiation of a  $vsx1^+$  progenitor at mitosis is not arbitrary, but matches that of the post-mitotic  $vsx1^+$  BCs in its vicinity.

### Results

### Bipolar cell progenitor mitoses occur over an extended timeperiod and relocate to non-apical sites

In common with many parts of the developing vertebrate CNS, the retina begins as a pseudostratified neuroepithelium with spindleshaped progenitors that span its apico-basal extent and undergo interkinetic nuclear migration, an oscillatory nuclear movement linked to specific cell cycle phases (Sauer, 1935; Baye & Link, 2008). At distinct but overlapping times, cells destined for different fates exit the cell cycle. Because mitotic divisions generally occur at the apical surface, newborn cells need to migrate varying distances to occupy their definitive locations within one of the emerging cellular laminae. Thus, while ganglion cells migrate furthest to occupy positions in the basal most part of the neuroepithelium, BCs have a shorter distance to relocate, and photoreceptors remain in situ at the apical surface. BCs, which are ultimately localized to the inner nuclear layer (INL) and confine their dendritic and axonal processes to the outer and inner plexiform layers (OPL, IPL), respectively, are generated over a protracted period, between 2 and 3 days postfertilization (dpf) in the zebrafish (He et al, 2012). Thus, early-born cohorts of BCs are generated when the retinal neuroepithelium is not yet laminated, while later-born cohorts are generated when the three cellular laminae are emerging.

To investigate the relationship between BC neurogenesis and differentiation, we examined the expression of *vsx1*, a transcription factor important for BC development (Passini *et al*, 1997; Chow *et al*, 2001; Vitorino *et al*, 2009; Shi *et al*, 2011), using a *vsx1*:GFP transgenic line (Kimura *et al*, 2008). In the zebrafish retina, *vsx1* is expressed at low levels in the majority of committed, terminally dividing BC progenitors, up-regulated during differentiation, and maintained at high levels in mature BCs (Vitorino *et al*, 2009). This developmental expression profile is faithfully reproduced in the bacterial artificial chromosome (BAC) *vsx1*:GFP transgenic line (Vitorino *et al*, 2009), so that we could follow nascent BCs from birth to maturity. Further, we took advantage of the zebrafish retina's gradient of development (Hu & Easter, 1999), to concurrently visualize immature (unlaminated) and more mature (laminated) parts of the retina and thus compare multiple states of BC differentiation in a single field of view.

At 2 dpf, we found low levels of GFP expression in the unlaminated part of the retina in which vsx1:GFP<sup>+</sup> (henceforth referred to as  $vsx1^+$ ) cells span the entire apico-basal axis. These  $vsx1^+$  cells represent both BC progenitors and post-mitotic, undifferentiated BCs (Morgan et al, 2006; Randlett et al, 2013). By contrast, vsx1<sup>+</sup> cells in the more mature, laminated part of the retina showed high levels of GFP expression and confined their processes to the IPL and OPL, suggestive of post-mitotic, differentiated BCs (Fig 1A). To our surprise, immunostaining of vsx1:GFP retinas for phosphorylated histone H3 (pH3), a late G<sub>2</sub>/M-phase marker (Hendzel et al, 1997), revealed  $vsx1^+$  pH3<sup>+</sup> cells not only in the unlaminated retina but also in the INL of the laminated retina. This suggests that not only post-mitotic BCs but also  $vsx1^+$  progenitors reside within the INL (Fig 1A–D). Notably, while  $vsx1^+$  progenitors in the unlaminated retina undergo mitosis at the apical surface, like "classical" progenitors in many parts of the CNS (Fig 1B), vsx1<sup>+</sup> progenitors in the laminated retina undergo mitosis in the INL (Fig 1C), akin to previously described non-apical progenitors (Godinho et al, 2007; Weber et al, 2014). Given the precocious expression of some neuronal characteristics in such non-apical progenitors, we asked how similar  $vsx1^+$  progenitors were to the post-mitotic  $vsx1^+$  BCs in their immediate surround.

### Marker expression in progenitors matches the surrounding postmitotic bipolar cells

GFP levels during mitotic division in  $vsx1^+$  pH3<sup>+</sup> progenitors showed a striking (3.8-fold) increase between the unlaminated and laminated regions of the developing retina (Fig 1E), which

# Figure 1. Vsx1<sup>+</sup> progenitors undergo mitosis in different proliferative zones and match the expression of molecular markers of post-mitotic BCs in their vicinity.

- A Confocal images of a coronal cryostat section from a 2 dpf usx1:GFP retina with immature, neuroepithelial ("unlaminated", orange) and mature, laminated regions ("laminated", cyan). Left panel, usx1:GFP; right panel, usx1:GFP shown in conjunction with pH3 antibody staining to label cells in late G<sub>2</sub>/M-phase. Scale bar: 10 µm.
- B, C High magnification images of (B) an apically dividing usx1<sup>+</sup> progenitor (orange arrowhead) in an unlaminated region where cells span the retina and express GFP weakly and (C) a non-apically dividing usx1<sup>+</sup> progenitor (cyan arrowhead) in a laminated region where cells confine their processes to the OPL and IPL (dashed lines) and express high levels of GFP. Cellular membranes are labeled with BODIPY methyl ester. Scale bar: 10 µm.
- D Quantification of  $vsx1^+$  progenitor mitoses at apical (lightly shaded) and non-apical (darkly shaded) locations in the unlaminated ("un", orange) and laminated ("lam", cyan) retina. Data are presented as mean  $\pm$  SEM, 1,391 mitotic divisions, 80 sections from at least 14 fish.
- E Quantification of usx1:GFP fluorescence intensity in dividing  $usx1^+$  progenitors in the unlaminated ("un", orange) and laminated ("lam", cyan) retina. Data are presented as mean  $\pm$  SEM, 86 progenitors, 43 sections from at least 13 eyes, Mann–Whitney U-test, \*\*\* $P \leq 0.0001$ .
- F Correlation of usx1:GFP fluorescence intensity in progenitors and their surrounding cells in the unlaminated (orange circles) and laminated (cyan circles) regions of the retina. Eighty-six progenitors from 43 sections from at least 13 eyes,  $r^2 = 0.78$ . Inset: Analysis for all pH3<sup>+</sup> cells (n = 13) from a single section.
- G Confocal images of a cryostat section from a 2 dpf usx1:GFP retina immunostained with antibodies against pH3 and Crx. Vsx1<sup>+</sup> progenitors (GFP<sup>+</sup> pH3<sup>+</sup>) are Crx negative (orange arrowheads) in the unlaminated retina (left panels, orange bars above panels) and are Crx positive (cyan arrowhead) in the laminated region (right panels, cyan bars above panels). Scale bar: 10 μm.
- H Correlation of Crx antibody staining intensity between mitotic BC progenitors and the  $usx1^+$  cells in their vicinity. Thirty-eight progenitors, 10 sections from at least five eyes,  $r^2 = 0.87$ ; orange circles, apical mitoses; cyan circles, non-apical mitoses.



Figure 1.

correlated well with the GFP levels in surrounding  $vsx1^+$  pH3<sup>-</sup> cells (Fig 1F; the surrounding  $vsx1^+$  cells in laminated regions are expected to be ~90% post-mitotic BCs on average; for an estimate of this number see Appendix Supplementary Materials and Methods). Furthermore, comparison of vsx1:GFP levels in pH3<sup>+</sup> progenitors and pH3<sup>-</sup> surrounding cells across the entire developmental gradient in single retinas revealed a linear increase along the gradient (Fig 1F, inset). Hence, with regard to vsx1 expression, progenitors in the laminated retina are more similar to their BC neighbors than to their early dividing peers and form a continuum with regard to vsx1 promoter activity in lock-step with surrounding BC differentiation. Direct time-lapse observation of vsx1:GFP levels confirmed a

parallel increase of fluorescence levels in BC progenitors and surrounding post-mitotic BCs *in vivo* (Fig EV1). Moreover, based on the decay of GFP in a *vsx2*:GFP BAC line (Vitorino *et al*, 2009), *vsx2* down-regulation was similarly linked to the progression of differentiation along the retinal gradient independent of mitotic status.

To further establish this similarity in molecular differentiation of  $vsx1^+$  progenitors and their post-mitotic neighbors, we examined two additional molecular markers of BC differentiation: cone-rod homeobox (Crx) and Ribeye a. Crx is a transcription factor expressed in mature photoreceptors and BCs (Liu *et al*, 2001; Shen & Raymond, 2004). In the unlaminated retina, antibody staining for



#### Figure 2. Morphological rearrangement of vsx1<sup>+</sup> progenitors matches surrounding BCs.

- A Confocal *in vivo* time-lapse recording of a retina from the Q26 transgenic line (crossed to a UAS:memYFP reporter) showing a non-apically dividing vsx1<sup>+</sup> progenitor (pseudo-colored cyan) with processes restricted to the IPL and OPL (dashed lines) during mitosis (0'). The last time point at which an apical process (open arrowhead) is detected is 89 min prior to mitosis. Scale bar: 10 µm.
- B Quantification of the distinct morphologies adopted by non-apically dividing usx1<sup>+</sup> progenitors at M-phase entry (122 progenitors, 17 fish). Open arrowheads indicate cytoplasmic processes extending beyond the synaptic layers (OPL and IPL, dashed lines).
- C Quantification of the time interval between retraction of the apical process (triangle) and mitosis (cyan circle) of non-apically dividing usx1<sup>+</sup> progenitors shows a broad range from 18 min to more than 540 min. As only mitosis, but not process retraction was observed for the progenitor depicted with small cyan dots, the movie length of 540 min is an underestimate. 18 progenitors from 11 fish.
- D Schematic of apical process retraction (triangle) in a non-apical usx1<sup>+</sup> progenitor (cyan soma) and the presence or absence of apical processes in the surrounding, post-mitotic BCs.
- E Quantification of the percentage of surrounding post-mitotic BCs without an apical process at the time when pre-mitotic  $usx1^+$  progenitors undergo apical process retraction (triangle, 60.3  $\pm$  6.6%) and at the time when these progenitors undergo mitosis (cyan circle, 95.1  $\pm$  1.7%). Data are presented as mean  $\pm$  SEM, 17 progenitors, 10 fish.

Crx (combined with pH3) revealed little or no expression in vsx1<sup>+</sup> progenitors or the  $vsx1^+$  cells in their vicinity, whereas in laminated regions, high levels of Crx expression were found in virtually all *vsx1*<sup>+</sup> progenitors and surrounding post-mitotic *vsx1*<sup>+</sup> BCs (Fig 1G). The levels of Crx antibody staining in mitotic vsx1<sup>+</sup> progenitors along the differentiation gradient strongly correlated with that of the *vsx1*<sup>+</sup> cells in their immediate surround (Fig 1H). Moreover, timelapse imaging of a crx:mCFP transgenic line (Suzuki et al, 2013), in which Crx<sup>+</sup> cells are faithfully labeled (Fig EV2A), regularly revealed non-apical crx:mCFP<sup>+</sup> mitotic divisions in the laminated retina (Fig EV2B and C). Finally, we analyzed the expression of Ribeye a, a structural protein of ribbon synapses in photoreceptors and BCs (Wan et al, 2005). As expected, using fluorescence in situ hybridization, we found ribeve a mRNA only in the laminated retina, where post-mitotic cells predominate (Fig EV2D and E). Notably, we also observed *ribeye a* mRNA-containing cells that were pH3<sup>+</sup> (Fig EV2F). The fact that these cells were located in the INL suggests they are BC progenitors. Using a transgenic line designed to report ribeye a expression in BCs (ctbp2:mEGFP; Odermatt et al, 2012), we observed *ctbp2*:mEGFP<sup>+</sup> cells dividing at non-apical locations, giving rise to BCs (Fig EV2G). Hence, even with regard to a marker linked to synaptic structures unique to BCs in the inner retina, we find that BC progenitors co-differentiate with post-mitotic BCs in the surround. We next asked whether this similarity extended beyond molecular markers to cellular morphology and dynamics.

# Progenitor morphology and cell biology correspond to the surrounding post-mitotic bipolar cells

To examine individual cells of the  $vsx1^+$  BC lineage, we generated a transgenic Gal4-driver line (referred to as Q26) that was selected to label a sparse subset of  $vsx1^+$  cells. As Gal4 expression in Q26 is restricted to  $vsx1^+$  BCs and their progenitors in laminated parts of the retinal gradient (Appendix Fig S1), we almost exclusively observed non-apically dividing Q26<sup>+</sup> progenitors, which we followed by time-lapse imaging as they terminally divided to produce BCs (Fig 2A). We examined the cleavage plane in dividing Q26<sup>+</sup> progenitors (n = 65) and found divisions along the circumferential, apico-basal, and centro-peripheral axes with no preference for a particular orientation, in line with other reports of nonapically dividing progenitors (Kimura *et al*, 2008; Weber *et al*, 2014). During mitosis, the vast majority of Q26<sup>+</sup> progenitors had limited their processes to the plexiform layers (Fig 2A at 0', Fig 2B), as is characteristic for BCs in the mouse and zebrafish retina (Morgan et al, 2006; Randlett et al, 2013) but not for apically dividing progenitors (Das et al, 2003; Miyata et al, 2004; Noctor et al, 2004). Prior to mitotic division however, the processes of non-apically dividing progenitors extended beyond the OPL or IPL (see Fig 2A at -89') and over time remodeled to become restricted to the synaptic layers. We asked whether this morphological remodeling occurred in a fixed time-window relative to mitotic division and focused our analyses, for technical reasons (see Appendix Supplementary Materials and Methods) on the retraction of processes from the apical surface. We found that apical process retraction occurred over an extended period of time (18 min to > 9 h) prior to mitotic division (Fig 2C). In contrast, for apically dividing BC progenitors in the unlaminated retina, apical process retraction to the OPL was a post-mitotic event (as observed in *vsx1*: GFP), but again occurred over an extended time span following mitotic division (a few min to > 8 h). Thus, a single differentiation step, the remodeling of the apical process, occurs both pre- and post-mitotically, and over a time span of more than 17 h relative to mitosis. Notably however, our time-lapse recordings in the Q26 line suggested that apical process remodeling is locally coordinated. When we identified progenitors that had just undergone apical process retraction to the OPL and asked whether post-mitotic BCs in the immediate vicinity had also done the same (Fig 2D), we found that, on the population level, apical process remodeling occurred concurrently (Fig 2E). Moreover, once pruned, the apical and basal processes of pre-mitotic progenitors could form lateral arbors. While these arbors regressed during mitosis, at earlier time points we could not distinguish them from the dendritic and axonal arbors of surrounding post-mitotic BCs (brackets in Fig 3A and E; Fig 4D at -645'), suggesting that the morphological processes of BC differentiation proceed independent of progenitor mitosis. In accordance with observations from other systems and species (Miyata et al, 2001; Das et al, 2003; Saito et al, 2003; Kosodo et al, 2008) the basally directed process of dividing  $Q26^+$  progenitors exhibited either splitting or asymmetric inheritance by one daughter cell followed by new outgrowth by the other daughter (Appendix Fig S2). Remarkably, when new process outgrowth was observed, it exhibited directed targeting of the IPL without overshooting beyond it.

Next, we asked whether  $vsx1^+$  progenitors undergo interkinetic nuclear migration. We used an mRNA construct encoding a fusion

#### Figure 3. Vsx1<sup>+</sup> progenitors in the unlaminated and laminated region exhibit distinct cell biological behaviors.

A Concurrent visualization of nucleokinesis in a *usx1*<sup>+</sup> progenitor in the laminated retina (*Q26*; *UAS*:memYFP, upper panels) and the cell cycle stage (mOrange2-PCNA mRNA, lower panels) by *in vivo* time-lapse confocal imaging. Progenitor soma is pseudo-colored cyan in YFP channel and outlined in cyan in mOrange channel. Scale bar: 10 μm.

B Nuclear movement of *usx1*<sup>+</sup> progenitors in the unlaminated region (*usx1*:GFP) of the retina prior to mitosis at the apical surface (orange circle). Seven cells, five fish. Black line represents the average position of all *usx1*<sup>+</sup> progenitors prior to mitosis in unlaminated regions.

C Nuclear trajectories of *vsx1*<sup>+</sup> progenitors in the laminated region that undergo mitosis at the INL/OPL interface (cyan circle, left panel) and post-mitotic BCs in their vicinity (right panel). In the time interval during which trajectories were tracked, all 12 *vsx1*<sup>+</sup> progenitors move to the INL/OPL interface prior to mitosis; 8 of 12 post-mitotic surrounding BCs do so (five fish). Black lines represent the average position of all *vsx1*<sup>+</sup> progenitors prior to mitosis in laminated regions (left panel) or of the post-mitotic BCs (right panel).

D Centrosomes (centrin4-YFP mRNA, grayscale) of a *usx1*<sup>+</sup> progenitor (magenta mask, *Q19*) and post-mitotic BCs in the laminated region concurrently translocate to the OPL and remain clustered there. The centrosome of the highlighted *usx1*<sup>+</sup> progenitor is pseudo-colored green. Scale bar: 10 μm.

E Continuation of the time-lapse from (D) (centrosomes not depicted). The  $vsx1^+$  progenitor (magenta mask) shows transient lateral arborizations in the synaptic layers as wide as its soma (see brackets at -210', -165'). A further example can be seen in (A) (see brackets at -67').

protein between mOrange2 and proliferating cell nuclear antigen (PCNA; Fig 3A and Appendix Fig S3) that allows determination of progenitor cell cycle phase by distinct nuclear localization patterns

(Leonhardt *et al*, 2000; Leung *et al*, 2011). Similar to classical progenitors,  $vsx1^+$  progenitors in the unlaminated region exhibited rapid apically directed interkinetic nuclear migration during G<sub>2</sub>







before mitosis at the apical surface (Fig 3B and Appendix Fig S3). This movement was absent in non-apically dividing  $vsx1^+$  progenitors, making their nuclear movements largely indistinguishable from the post-mitotic BCs in their immediate vicinity (Fig 3A and C).

However, the non-apically dividing  $vsx1^+$  progenitors always translocated their nuclei to the INL/OPL interface prior to mitosis. This movement can be explained by the location of the centrosome. By expressing fluorescently tagged centrin4, we found that BC

#### Figure 4. The differentiation status of late-born bipolar cells and earlier-born bipolar cells in their vicinity are similar.

A Fluorescence intensity of a non-apically dividing usx1<sup>+</sup> progenitor before it undergoes mitotic division (cyan trace), at the time of mitotic division (cyan circle) and of the two BC daughter cells (green ovals) over the course of 10 h (green traces).

- B Quantification of the fluorescence intensity of four pairs of BC daughters (green ovals) derived from non-apically dividing usx1<sup>+</sup> progenitors (cyan circle) 10 h after division. The fluorescence intensity of each of the BC pairs was normalized to the mitotic cell from which they were derived. BC pairs depicted are derived from two fish. The different overall rates of GFP fluorescence increase can be explained by where along the differentiation gradient cells originate.
- C Comparison of the *usx1* expression levels of apical progenitors, non-apical progenitors, and post-mitotic BCs derived from non-apically dividing progenitors 10 h following division. The fluorescence expression levels of cells surrounding each of the three cell categories are also quantified. Apically dividing *usx1*<sup>+</sup> progenitors (orange circle) and surrounding cells (dull orange bar, mean, and standard deviation, SD). Non-apically dividing *usx1*<sup>+</sup> progenitors (cyan circle) and surrounding cells (dull orange bar, mean, and standard deviation, SD). Non-apically dividing *usx1*<sup>+</sup> progenitors and 521 cells in their surround, and 43 non-apically dividing progenitors and 521 cells in their surround from at least 13 eyes were used for this analysis. Post-mitotic BCs (green oval) and earlier-born surrounding cells (dull green bar, mean, and standard deviation). Eleven post-mitotic BCs derived from seven mitotic divisions and 105 cells in their surround from three fish were used for this analysis.
- D Confocal *in vivo* time-lapse recording of the emergence of ribeye a puncta (pseudo-colored magenta, *ctbp2*:mCherry-ctbp2) in a BC derived from a non-apically dividing *vsx1*<sup>+</sup> progenitor (pseudo-colored cyan, *Q26*; *UAS*:memYFP), 7 h following cell cycle exit. Insets of the magenta-boxed region (at -118' and 0') of the axon terminal of the BC reveal the emergence of discernible ribeye a puncta at the 0' time point (white arrowheads). Scale bars: 10 µm.
- E Quantification of the time interval between mitosis and clustering of ribeye puncta (triangle) in the axon terminals of the ensuing BC daughters. Eight BCs derived from seven non-apically dividing progenitors (cyan circles) and eight BCs derived from apically dividing *vsx1*<sup>+</sup> progenitors (orange circles) from three fish were used for this analysis. In seven of the eight BCs derived from apical progenitors (depicted as small orange dots), we observed clustering but not the mitotic division, thus the time interval in these cases is an underestimate.
- F Quantification of the axon terminal arbor width of BCs derived from non-apical  $us \chi 1^+$  progenitors and the cells in their surround. At 10 h post-division, mean arbor width for BCs was 4.34  $\pm$  1.05  $\mu$ m, SD; the mean arbor width for cells in their surround was 4.29  $\pm$  1.24  $\mu$ m, SD. In total, nine BCs derived from seven non-apical divisions and 56 surround cells from six fish were used for analysis.

centrosomes actively relocated from the apical surface to the INL/ OPL interface (Fig 3D and E), resulting in a dendritic, rather than a somatic location at maturity (Fig EV3). Like apical process retraction, centrosome relocation occurred at the same time for progenitors and surrounding, post-mitotic BCs (Figs 3D and EV3). Thus, nucleokinesis and centrosome relocation represent further differentiation steps for which  $vsx1^+$  progenitors are time-locked to their post-mitotic neighbors.

### The post-mitotic differentiation status of late-born bipolar cells is similar to the early-born bipolar cell population in their vicinity

We assessed the rate of differentiation of the BC progeny resulting from non-apically dividing  $vsx1^+$  progenitors in the laminated region of the retina. First, we monitored vsx1-driven GFP expression levels in post-mitotic BCs over the course of 10 h after their exit from the cell cycle and found a steady increase in fluorescence intensity (Fig 4A). Ten hours post-division, GFP expression levels in BC sibling pairs were remarkably similar to each other (Fig 4A and B) and remained similar to the cells in their surround, which largely comprised of earlier-born post-mitotic BCs (Fig 4C).

Next, we used an established transgenic line, ctbp2:mCherryctbp2 (Pelassa et al, 2014), to examine the emergence of ribbon synapses in BC axon terminals. We assessed the time interval between exit from the cell cycle and the appearance of ribeye a clusters (marker of ribbon synapses) as an indication of presynaptic differentiation in vsx1<sup>+</sup> BCs (Fig 4D). For late-born BCs derived from non-apically dividing  $vsx1^+$  progenitors, the average time interval was 6.5 h. By contrast, for early-born BCs this interval was longer than 13 h on average (Fig 4E). Thus, late-born BCs acquire features of presynaptic differentiation with greater speed than their earlierborn counter-parts. Finally, we measured the lateral extent of the terminal axonal arbors of BCs derived from non-apical vsx1<sup>+</sup> progenitors 10 h after they exited the cell cycle, and found this to be remarkably similar to that of the earlier-born post-mitotic BCs in their vicinity (Fig 4F). Taken together, late-born BCs continue to differentiate in lock-step with earlier-born BCs in their vicinity, including the elaboration of markers of axonal and synaptic

differentiation, thus contributing to synchrony in local neuronal development.

# Experimentally delaying cell division does not delay usx1<sup>+</sup> progenitor differentiation

Our experiments have established that along the developmental gradient, BC progenitors blend into the differentiation landscape that surrounds them with regard to their morphological, cell biological, and molecular characteristics. Moreover, the "head-start" gained by pre-mitotic differentiation in late-dividing progenitors continues in their BC progeny, so that the lag in differentiation between late-born and early-born BCs is remarkably low. Two potential scenarios could explain these observations: (i) Multiple, "fixed" vsx1<sup>+</sup> progenitors exist, each of which undergoes mitosis at a stereotypic time point in the cell's differentiation trajectory. (ii) Alternatively, neurogenesis and differentiation could be uncoupled from each other, so that mitotic divisions could occur at various points in any given *vsx1*<sup>+</sup> cell's differentiation program. To distinguish between these two possibilities, we delayed BC progenitor divisions using hydroxyurea and aphidicolin (HUA). HUA treatment rapidly reduced the number of cells entering the G<sub>2</sub>/M-phase (Appendix Fig S4), but a small number of progenitors continued to divide, albeit with a prominent delay. We could now ask whether during delayed progenitor mitosis, differentiation stalled (as implied by the existence of "fixed" progenitors) or whether it continued and remained in synchrony with surrounding post-mitotic BC differentiation (as predicted by the "uncoupling" scenario; Fig 5A). We tracked 10 vsx1+ progenitors in HUA-treated embryos from late S-phase until mitosis. Knowing that progenitors, in control experiments, divided on average 142  $\pm$  3.9 min after the onset of late S-phase, we could determine when a HUA-treated cell should have divided ("expected" mitosis) and measured the delay with which the division actually occurred ("observed" mitosis, delay range approximately 3.5-9 h; Figs 5B and EV4). Because we could predict with a high degree of accuracy (86.4%), whether progenitors, in control conditions, would undergo mitosis at the apical surface or in the INL well before the divisions occurred (see



### Figure 5. Neurogenesis and differentiation of $\textit{vsx1}^+$ progenitors are independent of each other.

- A Schematic representation of expected outcomes if the immature retina (progenitors, P; neurons, N) is treated with HUA to delay the cell cycle. Upper panel: If there are multiple "fixed" progenitors, a block of cell division should stall progenitors at the differentiation state in which they normally would have undergone mitosis. The result would be a "salt-and-pepper" pattern of undifferentiated (light green) and differentiating progenitors (dark green). Lower panel: If cell cycle and differentiation are independent, all progenitors should homogenously differentiate. Open arrowheads indicate cytoplasmic processes not confined to the OPL and IPL, filled arrowheads indicate cytoplasmic processes confined to the synaptic layers.
- B Confocal images of a 2 dpf retina from a *usx1*:GFP embryo injected with a p53 morpholino and mOrange2-PCNA mRNA. A *usx1*<sup>+</sup> progenitor (dashed magenta outline) is shown before HUA treatment (left panel), at the time when it would have been "expected" to undergo mitosis (middle panel, orange diamond) and when it actually underwent mitosis (right panel, cyan circle). The retina and the *usx1*<sup>+</sup> progenitor continue to mature after the "expected" mitosis (retraction of cytoplasmic processes, mitosis at non-apical location, and up-regulation of GFP). Open arrowheads indicate cytoplasmic processes not confined to the OPL and IPL, filled arrowheads indicate cytoplasmic processes confined to the synaptic layers. Dotted lines indicate extent of *usx1*<sup>+</sup> cell somata across retinal thickness. Scale bar: 10 μm.
- C Progenitors that were expected to divide at the apical surface (exp), divided non-apically (obs). 10 progenitors, four fish.
- D Quantification of usx1:GFP fluorescence intensity of progenitors at the time when they were expected to undergo mitosis (exp, orange) and when they underwent mitotic division (obs, cyan). Data are presented as mean  $\pm$  SEM, 10 progenitors, four fish. Mann–Whitney *U*-test, \*\*\*P = 0.0002.
- E The fluorescence intensity of HUA-treated usx1:GFP progenitors at the time when they were expected to undergo mitosis (diamonds), and when they were observed to undergo mitosis (circles), plotted against the intensity of the surrounding cells (10 progenitors, four fish). One cell (cyan diamond) was expected to divide non-apically. Green line indicates the fluorescence change of the lowest expressing progenitor for clarity.

Appendix Supplementary Materials and Methods), we could ask whether the HUA-induced delay of mitosis would shift divisions from apical to non-apical locations (as an "uncoupling" scenario would imply). Nine of the 10 HUA-treated cells fulfilled criteria that identified them to be destined to divide apically. However, all 10 cells instead underwent mitosis in the INL after process remodeling, suggesting that they had been shifted from an apical to a non-apical phenotype simply by delaying mitosis (Fig 5C). Furthermore, vsx1: GFP expression levels in the 10 delayed progenitors increased from the point of expected mitosis to observed mitosis (Fig 5D). Notably, even hours after the delayed mitosis occurred, progenitors still matched the fluorescence levels of cells in their immediate surround (Fig 5E), suggesting uncoupling of cell division and differentiation. Together, these findings support the "uncoupling" scenario laid out above and hence argue for an independence of neurogenesis and differentiation programs during BC development.

## Discussion

Our study is the first, to our knowledge, to address the relative timing of neurogenesis, migration, and differentiation for a molecularly defined CNS population in vivo and to elucidate the effects of such timing on progenitor characteristics. In contrast to the widely held view, we found that developing neurons did not adopt a stereotypic sequence of neurogenesis followed by migration and subsequent differentiation. Rather, to our surprise, the developmental trajectories that progenitors adopted were variable and accommodated remarkable flexibility. This resulted in vsx1<sup>+</sup> BC progenitors with a wide variety of molecular, morphological, and cell biological characteristics. Importantly, rather than representing many distinct populations, the differentiation status of these progenitors formed a continuum in lock-step with the differentiation of surrounding postmitotic BCs along the developmental gradient of the retina. Accordingly, cells dividing in the laminated parts of the retina were more BC-like than their early dividing counter-parts. Indeed, without the aid of time-lapse imaging or cell cycle markers it would have been impossible to distinguish between pre-mitotic progenitors and postmitotic cells. Our results support the conclusion that a stereotypical and fixed sequence of ontogenetic events is not essential during neuronal development. Thus, at least for terminally dividing progenitors, which generate a substantial part of the CNS neuronal population (Nakashima et al, 2015), mitosis does not have to occur before neuronal differentiation is initiated (or at any specific step thereafter), but rather can be flexibly intercalated between other developmental steps.

Our findings could explain two previous intriguing observations: First, that neuronal progenitors in different parts of the nervous system show precocious signs of differentiation prior to cell cycle exit (Rothman *et al*, 1980; Rohrer & Thoenen, 1987; DiCicco-Bloom *et al*, 1990; Miyata *et al*, 2004; Godinho *et al*, 2007; Attardo *et al*, 2008; Prasov & Glaser, 2012); and second, that blocking mitosis does not halt neuronal differentiation in many parts of the *Xenopus* CNS including the retina, spinal cord, and brain stem (Harris & Hartenstein, 1991). Our results now offer a unifying explanation for these previous observations, suggesting that they might simply result from a fundamental uncoupling of cell cycle and neuronal differentiation during normal development. Indeed, the increasingly recognized prevalence of precocious progenitors in many parts of the nervous system of a range of species [e.g., basal progenitors in the neocortex (Haubensak *et al*, 2004; Miyata *et al*, 2004; Noctor *et al*, 2004); neural crest-derived PNS progenitors (Rothman *et al*, 1980; Rohrer & Thoenen, 1987; DiCicco-Bloom *et al*, 1990); retinal progenitors (Godinho *et al*, 2007; Prasov & Glaser, 2012)], implies that such uncoupling may be a general principle of neural development.

What would be the advantages of uncoupling neurogenesis and neuronal differentiation? By comparison with the orderly sequence of developmental events in the classical model, we suggest that the uncoupled model provides two advantages, namely speed and synchrony (Fig EV5). In the classical model, because differentiation can only be initiated following mitosis, the time required for the majority of cells in a defined population to differentiate is dictated by the delay in their birth dates. By contrast, in the uncoupled model, differentiation steps can already occur at the progenitor stage, permitting differentiation across the population to be faster. Furthermore, despite the extended time span over which mitotic divisions can occur, cells in a given population differentiate in relative synchrony. The uncoupling of mitosis and differentiation could thus be particularly pertinent for the assembly of essential neural circuits where swift maturation is paramount for survival (Nikolaou & Meyer, 2015).

## Materials and Methods

### Animals

All experiments were performed according to regulations as approved by the local regulatory bodies. Zebrafish were maintained, mated, and raised as described in Mullins *et al* (1994). Embryos were kept in  $0.3 \times$  Danieau's solution at  $28.5^{\circ}$ C and staged as previously described (Kimmel *et al*, 1995). Fish were either in an AB wild-type or *roy orbison* (Ren *et al*, 2002) background. The transgenic lines used are listed in Appendix Table S1. We generated Tg (*vsx1*:Gal4)q26 (*Q26*) and Tg(14*xUAS*:memTagRFP-T) by Tol2 mediated insertion (Kawakami, 2004). For details of constructs used to make transgenic fish and for transient injections see Appendix Supplementary Materials and Methods.

### mRNA synthesis and injection

Plasmids were linearized (PCNA: NotI, centrin4: ApaI). Capped mRNA was produced using the Ambion mMESSAGE mMACHINE kit (Applied Biosystems) according to the manufacturer's instructions. mRNA was injected at 100 ng/µl into one- or two-cell stage embryos.

#### Immunohistochemistry and in situ hybridization

Immunostaining to detect pH3, Crx, and GFP was performed using adaptations of previously published protocols either on cryosections (Williams *et al*, 2010) or on whole-mount embryos (Hunter *et al*, 2011). *In situ* hybridization to visualize *ribeye a* mRNA was performed on whole-mount embryos using a digoxigenin-labeled riboprobe and Fast Red TR/naphthol AS-MX (Sigma) to detect alkaline phosphatase activity. For a detailed description see Appendix Supplementary Materials and Methods.

### In vivo imaging

Embryos were prepared for imaging as described previously (Godinho, 2011; Engerer *et al*, 2016). Between 10 and 18 h post-fertilization (hpf), embryos were transferred to  $0.3 \times$  Danieau's solution containing 0.003% 1-phenyl-2-thiourea (PTU, Sigma) to inhibit melanin formation. At 2.25 days post-fertilization (dpf), manually dechorionated embryos were anesthetized using 0.02% tricaine (PharmaQ) in medium containing PTU and embedded laying on their side in low-melting agarose (0.7–0.8%, Sigma).

Fish were imaged starting at 2 dpf on an Olympus FV1000 confocal/2-photon and an Olympus FVMPE-RS 2-photon microscope using water-immersion objectives (Olympus  $20\times/NA 0.95$ , Olympus  $25\times/NA 1.05$ , Zeiss  $40\times/NA 1.0$ , Nikon  $25\times/NA 1.1$ , and Nikon  $40\times/NA 0.8$ ) or a silicon-immersion objective (Olympus  $30\times/NA$ 1.05). Embryos were maintained at 28.5°C during all *in vivo* recordings. At each time point *z*-stacks were acquired of the peripheral retina, encompassing its entire circumference.

### Image processing

Images were viewed and processed using open-source ImageJ/Fiji software (http://fiji.sc). The StackReg function was used for drift compensation in xy. Image panels were assembled in Photoshop CS5 (Adobe) and combined into figures using Illustrator CS5 (Adobe). The "Gaussian blur" function was used to filter noise for clarity. Gamma was not adjusted.

### Hydroxyurea-aphidicolin treatment

Hydroxyurea (Sigma) and aphidicolin (BioViotica) were used at a final concentration of 20 mM and 150  $\mu$ M, respectively, in 0.3× Danieau's containing 1.0–1.7% DMSO. Embryos were injected with a p53 morpholino (0.5–1 mM, Gene Tools) at the one- or two-cell stage to ameliorate HUA-induced apoptosis (Girdler *et al*, 2013). At 2 dpf embryos were mounted in agarose as described for *in vivo* imaging above, but leaving the tail fin un-embedded for better drug access. Retinas were imaged for one time point prior to HUA administration. The 0.3× Danieau's medium was replaced with HUA-containing medium, and time-lapse recording was immediately resumed. Recordings were generally limited to < 16 h after HUA addition as high levels of cell death were observed thereafter.

### Data analysis

For details, see Appendix Supplementary Materials and Methods.

### Statistics

Mean values and standard error of the mean (SEM) were calculated using Microsoft Excel. We used the Mann–Whitney *U*-test to compare datasets with GraphPad Prism 5. Data are presented as mean  $\pm$  SEM unless indicated otherwise. *P*-values < 0.05 are denoted with "\*", *P* < 0.01 with "\*\*", and *P* < 0.001 with "\*\*\*".

Expanded View for this article is available online.

### Acknowledgements

We thank Kristina Wullimann for expert fish husbandry and Yvonne Hufnagel. Alexandra Graupner, and Monika Schetterer for expert technical and administrative support. We are especially grateful to Rachel Wong (U. Washington, Seattle), who generously supported this project in multiple ways, thanks to her, and Magdalena Götz and Tim Czopka for critical reading of an earlier version of this manuscript. We are grateful to M. Nonet (Washington University in St. Louis) for the pCold Heart Tol2 vector, M. Meyer (King's College London) for the 5xUAS:TagRFP-T vector, T. Nicolson (Oregon Health & Science University and Vollum Institute) for the ribeye a cDNA template, R. Köster (Technische Universität Braunschweig) for pSK5xUAS:Centrin2-YFP, and P. Raymond (University of Michigan) for the anti-Crx antibody. The mOrange2-PCNA-19-SV40NLS-4 plasmid was a gift from Michael Davidson (Addgene plasmid # 57971). We thank S. Higashijima (National Institutes of Natural Sciences, Okazaki Institute for Integrative Bioscience) for the Tg(vsx1:GFP)nns5 and Tg(vsx2:GFP)nns1 BAC transgenic lines which were generated with the support of the National Bioresource Project of Japan. We also thank R. Wong for providing the following transgenic lines: Tg(pax6-DF4:gap43-CFP)q01, Tg (vsx1:MCerulean)q19, Tg(14xUAS:MYFP), and Tg(crx:MA-CFP)q20, and L. Lagnado (University of Sussex, UK) for Tg(-1.8ctbp2a:mCherry-ctbp2a)Imb7. The Tg(-1.8ctbp2:gap43-EGFP)Imb1 line was made in the laboratory of L. Lagnado at MRC-LMB, Cambridge. This project was made possible by funding to L.G. and T.M. from the Deutsche Forschungsgemeinschaft (DFG) through Collaborative Research Center 870 "Assembly and Function of Neuronal Circuits", project A11. T.M. is further supported by the Center for Integrated Protein Science Munich (CIPSM, EXC 114), the European Research Council under the European Union's Seventh Framework Program (FP/2007-2013; ERC Grant Agreement no. 616791), the Munich Center for Systems Neurology (SyNergy; EXC 1010), the DFG Priority Program 1710 (Mi694/4-1), the DFG research grants Mi694/7-1, 8-1, and the German-Israeli Foundation (I-1200-237.1/2012). T.M. is also associated with the German Center for Neurodegenerative Diseases (DZNE Munich). P.R.W. was supported by the Human Frontier Science Program and the Wings for Life Foundation. N.O. was supported by the Amgen Scholar program. S.C.S. and T.Y. were supported by a grant awarded to R. Wong (NIH EY14358). P.E. was supported by the DFG Research Training Group 1373 and the Graduate School of the Technische Universität München (TUM-GS).

### Author contributions

PE, PRW, TM, and LG conceived of the project and designed experiments. TM and LG supervised the project. PE performed all *in vivo* imaging experiments, with some help from NO. PE and PC performed *in situ* hybridization. PE, SCS, TY, NO, and BO generated new constructs and transgenic lines for this research. PE, PRW, TM, and LG wrote the paper with input from all authors.

### **Conflict of interest**

The authors declare that they have no conflict of interest.

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