Notch-mediated re-specification of neuronal identity during central nervous system development

Highlights

- Spontaneous fate reprogramming in the vsx1 lineage of the zebrafish retina
- Nascent vsx1 bipolar cells are re-specified to an amacrine cell fate
- Notch signaling plays a major role in vsx1 bipolar cell respecification
- Over-stimulating Notch signaling enhances reprogramming in the vsx1 lineage

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In brief

Engerer, Petridou et al. find evidence for spontaneous cell fate reprogramming during retinal development. Newly postmitotic *vsx1* bipolar cells switch to an amacrine cell fate. Notch signaling plays a major role, conferring plasticity to nascent bipolar cells, allowing suitable transcription factors to instruct the amacrine cell fate.









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Notch-mediated re-specification of neuronal identity during central nervous system development

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SUMMARY

Neuronal identity has long been thought of as immutable, so that once a cell acquires a specific fate, it is maintained for life.¹ Studies using the overexpression of potent transcription factors to experimentally reprogram neuronal fate in the mouse neocortex²,³ and retina⁴,⁵ have challenged this notion by revealing that post-mitotic neurons can switch their identity. Whether fate reprogramming is part of normal development in the central nervous system (CNS) is unclear. While there are some reports of physiological cell fate reprogramming in invertebrates,⁶,² and in the vertebrate peripheral nervous system,⁶ endogenous fate reprogramming in the vertebrate CNS has not been documented. Here, we demonstrate spontaneous fate re-specification in an interneuron lineage in the zebrafish retina. We show that the *visual system homeobox 1* (*vsx1*)-expressing lineage, which has been associated exclusively with excitatory bipolar cell (BC) interneurons,⁶¹²² also generates inhibitory amacrine cells (ACs). We identify a role for Notch signaling in conferring plasticity to nascent *vsx1* BCs, allowing suitable transcription factor programs to re-specify them to an AC fate. Overstimulating Notch signaling enhances this physiological phenotype so that both daughters of a *vsx1* progenitor differentiate into ACs and partially differentiated *vsx1* BCs can be converted into ACs. Furthermore, this physiological re-specification can be mimicked to allow experimental induction of an entirely distinct fate, that of retinal projection neurons, from the *vsx1* lineage. Our observations reveal unanticipated plasticity of cell fate during retinal development.

RESULTS AND DISCUSSION

Vsx1 progenitors generate bipolar cells (BCs) and amacrine cells (ACs)

Vsx1 is a key determinant of BC fate across vertebrate species. 9-11,13 In the zebrafish retina, *vsx1* BCs are generated by *vsx1* progenitors via terminal, symmetric mitotic divisions, 14,15 suggesting they are hard-wired to the BC fate. However, in a *vsx1*:GFP transgenic line that faithfully reports *vsx1* expression, 12,16 small numbers of ACs are labeled (Figure 1A). We investigated the origin of these ACs by *in vivo* time-lapse recordings in a Gal4-driver line (*Q32*) that provides a lineage label for a

subset of vsx1 cells.¹⁴ ACs were generated by terminally dividing Q32 progenitors, with BCs as their siblings. These AC-BC divisions represented almost half of all Q32 divisions (41.2% \pm 6%, mean \pm SEM, 141 divisions, 16 fish; Figure 1B; see also lineage tracing studies^{17,18}). Q32 daughters that became ACs maintained somal contact with their BC sibling for several hours (5.3 \pm 0.7 h, mean \pm SEM, 13 pairs, 6 fish), before acquiring AC features. By contrast, BC-BC siblings did not lose somal contact after mitosis (at least over the recording, 14.5 ± 0.7 h, mean \pm SEM, 10 pairs, 7 fish). Loss of somal contact with their BC sibling thus represents the earliest sign of morphological differentiation toward an AC fate (Figure S1A). Q32 ACs express a bona fide pan AC marker. $ptf1a^{19}$



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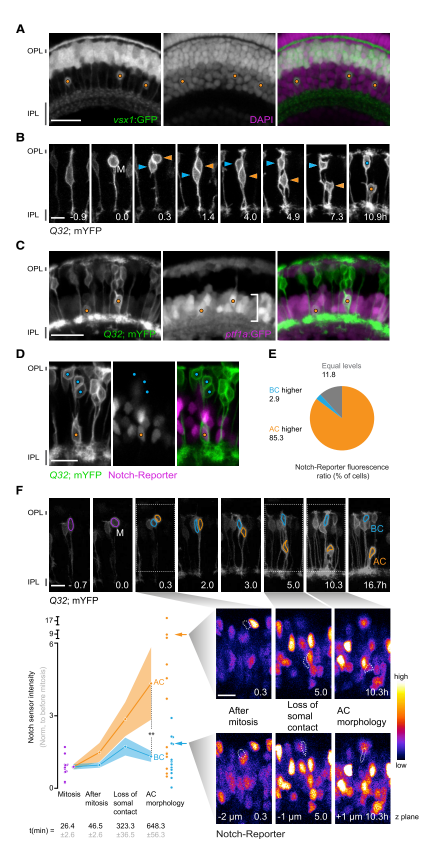


Figure 1. Vsx1 progenitors generate ACs

(A) 3 dpf vsx1:GFP retina with vsx1+ BCs and ACs (orange circles).

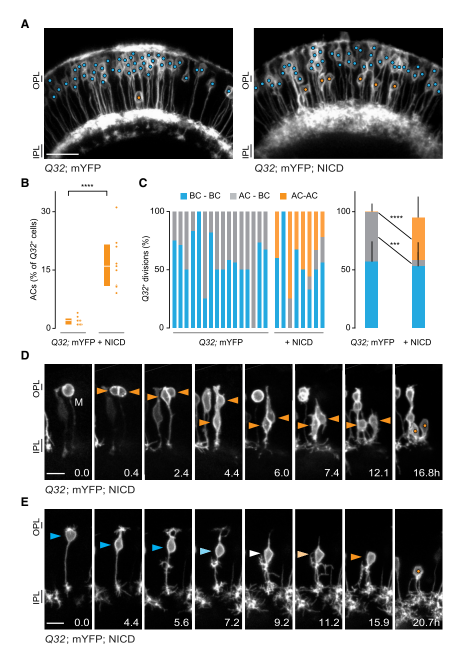
(B) 2 dpf Q32 progenitor undergoing mitosis (M), generates a BC (cyan arrowhead; circle, final time point) and an AC (orange arrow-head, circle final time-point).

(C) 3 dpf WT retina showing ptfa:GFP+ Q32 ACs (orange circles). Ptf1a:GFP labels all ACs (INL, bracket, center panel). Ptf1a-GFP signal bleeds through Q32-YFP channel.

- (D) 3 dpf retina showing Notch-reporter (tp1:hmgb1mCh erry) expression in a Q32 AC (orange circle), but not in surrounding Q32 s BCs (cyan circles).
- (E) Notch-reporter (tp1:hmgb1mCherry) fluorescence intensity in Q32 ACs and BCs. AC and BC categories were tested against an expected frequency of 50% using a binomial test, p < 0.0001; 30 cells, 10 fish. Four ACs (of 34) displayed equal levels of fluorescence as neighboring BCs.
- (F) 2 dpf transgenic retina (Q32; mYFP; tp1:H2BmCherry; top grayscale images, gamma adjusted) showing a WT Q32 progenitor (purple outline) generating a BC (cyan) and an AC (orange). Notch-reporter levels in the AC and BC depicted using a Fire LUT (below, right). Notch-reporter intensity levels (means ± SEMs) for 13 Q32 progenitors (8 fish) and their BC (cyan) and AC (orange) daughters (below, left). Times in relation to the time point before mitosis. Significant differences between BC-AC pairs were found when the AC acquired its characteristic morphology (Wilcoxon matched-pairs signed-rank test,

Scale bars, 20 μm (A and C); 10 μm (B, D, and F). IPL, inner plexiform layer; OPL, outer plexiform layer. See also Figures S1A, S2, and S3A-S3C.





(99.2% \pm 0.8% of Q32 ACs were ptf1a:GFP+; mean \pm SEM, 60 cells, 13 fish; Figure 1C), and only a small number maintain vsx1 expression (1 of 26 Q32 ACs was vsx1:GFP+ at 3 days post-fertilization [dpf], 15 fish; Figures S2A–S2C), accounting for the low AC number in vsx1:GFP retinae (0.8% \pm 0.2% of 2,295 vsx1:GFP+ cells, mean \pm SEM, 3 dpf, 8 fish). We also detected cells with an AC morphology in BC transgenic lines (crx:mCFP and ctbp2:mEGFP; 14,20,21 Figure S2D), hinting at their BC lineage origin.

Notch signaling promotes the AC fate in the *vsx1* lineage

To investigate whether Notch signaling underlies the asymmetric fates, ^{16,22–26} we crossed the *Q32* line to Notch-reporter lines^{27,28} (Figure 1D). The majority of *Q32* ACs displayed higher Notch-

Figure 2. Notch signaling promotes AC fate in the *vsx1* lineage

(A) 3 dpf retinae from Q32 WT and Q32 NICD. BCs (cyan circles), ACs (orange circles).

(B) Proportion of ACs in WT and Q32 NICD. Median and interquartile range (IQR) shown. Q32 WT (21 ACs, 9 fish); Q32 NICD (140 ACs, 9 fish), p < 0.0001, Mann-Whitney U test.

(C) Proportion of Q32 divisions in WT and following NICD OE for each fish (left). Collective representation of Q32 divisions, representing median and IQR per group (right). BC-BC: Q32 WT versus NICD, p = 0.5134. AC-BC: Q32 WT versus NICD, p = 0.0004. AC-AC: Q32 WT versus NICD, p < 0.0001. WT: 141 Q32 divisions, 16 fish; Q32 NICD: 44 divisions, 8 fish. Mann-Whitney U test.

(D) 2 dpf Q32 NICD retina, showing a mitotic progenitor (M) generating 2 ACs (orange arrowheads; circles, last time point).

(E) 2 dpf Q32 NICD retina, showing a BC (cyan arrowhead), transdifferentiating into an AC (orange arrowhead; circle, last time point).

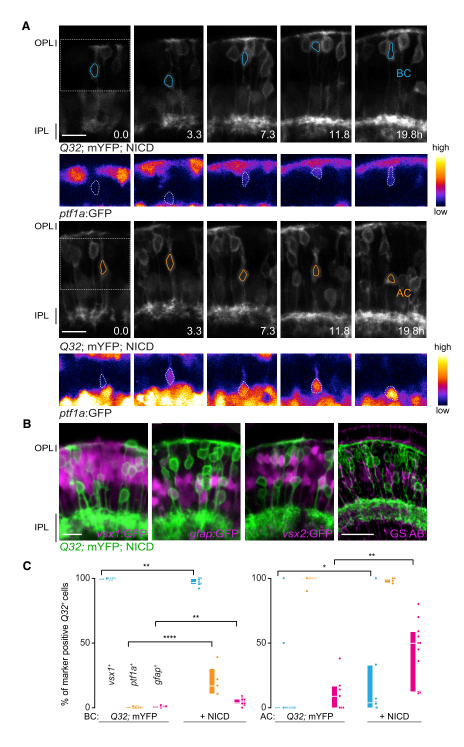
Scale bars, 20 μ m (A); 10 μ m (D and E). See also Figures S1, S3D, and S3E.

reporter levels than Q32 BCs in their vicinity (Figure 1E). To determine when the divergence of Notch signaling emerged, we monitored Notch-reporter expression in time-lapse recordings (Figure 1F). Notchreporter levels were low in Q32 progenitors and increased gradually post-mitosis. Significant increases were only detectable in the AC sibling of AC-BC pairs when it acquired its typical morphology (average 10 h post-mitosis), implicating the postmitotic nature of this signaling event. We investigated whether mechanisms at the progenitor level²⁹⁻³¹ accounted for the signaling asymmetry in the ensuing daughters. However, we found neither asymmetric distribution of the Notch regulator Numb (10 cells, 4 fish) nor mitotic cleavage along a stereotypic axis (45 cells, 25 fish; Figures S3A and S3B), suggesting these key players are not involved.

We used DAPT, a γ -secretase inhibitor, to abrogate Notch signaling and found an \sim 50% decrease in vsx1:GFP+ ACs (DAPT-treated 2.0 \pm 1.4 ACs per region, 23 regions, 12 fish versus DMSO controls 5.0 \pm 1.6 ACs per region, 18 regions, 9 fish, median \pm SD, p < 0.0001, Mann-Whitney U test; Figure S3B). Conversely, overexpression (OE) of the Notch intracellular domain (NICD) in the vsx1 lineage (Q32 NICD) led to a 9-fold increase in Q32 ACs (Figures 2A and 2B), that persisted at least until 5 dpf (Figure S3C). The increased AC number did not arise from increased Q32 progenitor proliferation (0.21 \pm 0.03 wild type [WT] versus 0.19 \pm 0.05 NICD, means \pm SEMs, divisions per hour, monitored 18 h from 2 to 3 dpf; p = 0.75, t test), but rather from the emergence of AC-AC pairs, a division mode not seen in Q32 WT (Figures 2C and 2D), and by BC-to-AC

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transdifferentiation (17 observations, 9 fish; Figure 2E). Transdifferentiating BCs exhibited process exuberance and took longer to acquire AC features compared to WT (Figure S1B), highlighting the fate switch that occurred along their original differentiation trajectory. Time-lapse of the Notch-reporter in Q32 NICD during transdifferentiation showed that Notch signaling increases when the soma translocated toward the inner part of the inner nuclear layer (INL) and AC morphology emerged. Following AC-BC divisions, Notch-reporter levels increased Figure 3. Marker expression in re-specified

(A) In Q32 NICD, BCs (outlined cyan, upper grayscale panels), express ptf1a:GFP (Fire LUT), albeit at lower levels than in ACs (outlined orange, lower grayscale panels). Ptf1a-GFP signal bleeds through Q32-YFP channel (grayscale panels). (B) Following NICD OE, transdifferentiated Q32

ACs do not express vsx1:GFP but gfap:GFP. Q32 ACs are negative for vsx2:GFP and glutamine synthetase (GS antibody, AB).

(C) Percentage of Q32 BCs (left) and ACs (right) expressing vsx1, ptf1a and gfap in WT or Q32 NICD. Median and IQR represented. BCs: vsx1:GFP+: Q32 WT, (1227 BCs, 14 fish) versus NICD, (416 BCs, 7 fish); p = 0.0034. Ptf1a:GFP+: Q32 WT (1,614 BCs, 14 fish) versus NICD (486 BCs, 5 fish); p < 0.0001. Gfap:GFP+: Q32 WT (718 BCs, 7 fish) versus NICD (984 BCs, 11 fish); p = 0.0022. ACs: vsx1:GFP+: Q32 WT (28 ACs, 15 fish) versus Q32 NICD (100 ACs, 7 fish); p = 0.0129. Ptf1a:GFP+: Q32 WT (60 ACs, 13 fish) versus Q32 NICD (124 ACs, 4 fish); p = 0.1206. Gfap:GFP+: Q32 WT (43 ACs, 7 fish) versus Q32 NICD (335 ACs, 11 fish); p = 0.0078. Mann-Whitney U test for pairwise comparisons.

Scale bars, 10 µm (A and B).

significantly in the AC sibling several hours (average 6 h) post-mitosis (Figures S1C and S1D). Thus, also after NICD OE, Notch signaling appears to operate post-mitotically to re-specify fate.

Transdifferentiated ACs lose BC markers and express markers of **immaturity**

ACs in Q32 NICD retinae were ptf1a:GFP+ and most lost expression of vsx1:GFP (Figures 3A and 3C). The NICD-induced ACs also lacked other BC markers (0% crx:mCFP+, 65 ACs, 3 fish; 1.9% ± 1.1% ctbp2:mEGFP+, means ± SEMs, 95 ACs, 4 fish), resembling WT Q32 ACs. Notably, 20% of BCs in Q32 NICD expressed ptf1a:GFP, a phenotype that is absent in WT fish (Figures 3A and 3C). This suggests that after NICD OE, some BCs expressed AC markers, but did not adopt an AC morphology. Moreover, >40% of Q32 NICD ACs were gfap:GFP+, which

is expressed in neural progenitors^{34,35} (Figures 3B and 3C). Importantly, ~11% of WT Q32 ACs also expressed this marker. Notch signaling is associated with Müller cell specification^{36–39} and GFAP is expressed by Müller glia.³⁴ To exclude that NICDinduced Q32 ACs were driven toward a Müller cell fate, we examined their expression of glutamine synthetase (GS)⁴⁰ and vsx2, 12 but found no evidence for this (GS: 0 of 68 ACs, 7 Q32 NICD fish; 0 of 28 ACs, 9 Q32 WT fish; vsx2:GFP: 0 of 274 ACs, 6 Q32 NICD fish; 0 of 26 ACs, 6 Q32 WT fish). Thus, gfap



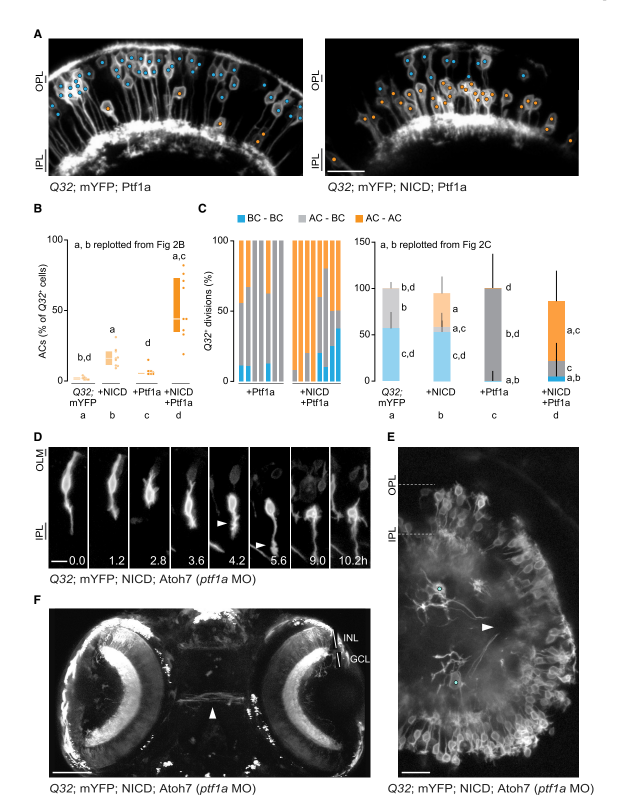


Figure 4. Notch signaling imparts plasticity to nascent post-mitotic cells (A) 3 dpf retinae from Q32 Ptf1a and Q32 NICD Ptf1a fish. Q32 BCs (cyan circles), ACs (orange circles). (B) Proportion of ACs in Q32 Ptf1a and Q32 NICD Ptf1a represented as median and IQR. Q32 Ptf1a (66 ACs, 9 fish), Q32 NICD Ptf1a (372 ACs, 9 fish). Q32 WT and Q32 NICD originally plotted in Figure 2B. Significant differences found between WT and NICD (p = 0.0003), WT and NICD Ptf1a (p < 0.0001), and Ptf1a and NICD

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in Q32 ACs does not indicate Müller cell identity, but rather an immature state, permissive of a fate switch.

Ptf1a can induce the AC fate in the vsx1 lineage

We next asked whether Ptf1a, a determinant of AC fate, 41-45 also governs the fate of vsx1 ACs. Ptf1a OE in the vsx1 lineage (Q32 Ptf1a) resulted in a 3.6-fold increase in Q32 ACs compared to WT (Figures 4A and 4B). Divisions generating at least 1 AC were higher in Q32 Ptf1a (100% \pm 6.19%, median \pm SD) than in Q32 WT (43% ± 23.8%, median ± SD, Mann-Whitney U test, p = 0.0001; Figure 4C) and in Q32 NICD (47% \pm 28.5%, median \pm SD, Mann-Whitney U test, p = 0.0048). Surprisingly, we found a lower number of excess ACs in Q32 Ptf1a compared to Q32 NICD (Figure 4B), likely due to the lack of transdifferentiation events. Thus, Ptf1a appears to influence fate around mitosis or shortly thereafter. To examine whether NICD OE could extend this plasticity window, we generated Q32 NICD Ptf1a fish and found a 28-fold increase in Q32 ACs compared to WT (Figures 4A and 4B). AC-AC pairs accounted for the majority of divisions in these fish (66.5%; Figure 4C). Thus, physiological Notch signaling confers plasticity to nascent vsx1 BCs, making them receptive to Ptf1a, which instructs the switch to an AC fate, a phenomenon that can be enhanced by combined NICD and Ptf1a OE. We next probed whether NICD OE combined with another transcription factor could instruct a cell fate beyond ACs.

Ganglion cell (GC)-like cells can be induced in the vsx1 lineage

We tested whether GCs could be generated by knocking down ptf1a43,46 to suppress AC fate and overexpressing NICD with atonal homolog 7 (Atoh7), a critical GC fate determinant. 47-51 We observed bipolar-shaped Q32 cells, transforming morphologically while translocating to the GC layer (GCL) and initiating axon outgrowth (14 cells, 3 fish; Figure 4D). These cells expressed isl2b:GFP, a GC marker⁵² (Figure S4). Of 68 Q32 GCL-localized cells (24 fish), 30 bore morphologies reminiscent of GCs (Figure 4E), while the rest lacked a detectable axon or displayed few or no discernible neurites. At 5 dpf (4 fish), we observed Q32 GC axons at the optic chiasm, suggesting successful retinal exit and path finding en route to the optic tectum (Figures 4F and S4).

Implications of developmental plasticity

Here, we report physiological Notch-dependent fate reprogramming in the vsx1 lineage, allowing nascent BCs to re-specify to an AC fate. That Notch signaling confers plasticity to nascent postmitotic cells has been suggested to occur in the postnatal mouse retina.⁵³ Notch signaling has also been implicated in transdifferentiation events in the damaged mammalian cochlea.⁵⁴ In the latter case, however, in contrast to our findings, attenuation of Notch signaling is involved. 55

Vsx1-lineage-derived ACs represent a small fraction of the total AC population, the majority of which arise from a different lineage. 43,56 Could vsx1 ACs have a specific functional significance? One can speculate that vsx1 ACs are generated "on demand," with the BC-to-AC re-specification allowing local fine-tuning of excitation and inhibition. Reducing the AC number, using a ptf1a morpholino, led to an almost 2-fold increase in vsx1 cells expressing ptf1a:GFP, thus commencing differentiation toward an inhibitory fate.

Finally, understanding the mechanisms of physiological fate re-specification could guide cell replacement strategies such as direct *in vivo* reprogramming.⁵⁷ While progress has been made converting glia to neurons, ^{58,59} re-specifying neurons from one subtype to another is limited to a short developmental time window, perhaps due to epigenetic changes that fixate cell type identities.^{2,3,5} A pulse of Notch signaling may rejuvenate post-mitotic cells and re-open the window for transcription factor-induced re-specification of neuronal subtypes and thus should be explored in the ongoing search for efficient replacement strategies.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

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Ptf1a (p = 0.0005), denoted by a (significantly different from Q32 WT), b (from NICD), c (from Ptf1a), or d (from NICD Ptf1a). Kruskal-Wallis and post hoc Dunn's multiple comparisons test, further adjusted by the Benjamini-Hochberg false discovery rate (FDR) method.

(C) Q32 divisions in Q32 Ptf1a and Q32 Ptf1a NICD for each fish (left). Divisions representing median and IQR per group (right). Q32 Ptf1a: 42 divisions, 7 fish; Q32 NICD Ptf1a: 52 divisions, 8 fish. Q32 WT and Q32 NICD originally plotted in Figure 2C. Significant differences were found in BC-BC divisions: WT versus Ptf1a (p = 0.0001), WT versus NICD Ptf1a (p = 0.0004), NICD versus Ptf1a (p = 0.0029), NICD versus NICD Ptf1a (p = 0.0075); AC-BC divisions: WT versus NICD (p = 0.0127), NICD versus Ptf1a (p = 0.0001), Ptf1a versus NICD Ptf1a (p = 0.0034); AC-AC divisions: WT versus NICD (p = 0.0007), WT versus NICD Ptf1a (p < 0.0001), Ptf1a versus NICD Ptf1a (p = 0.0035), denoted as in (B). Kruskal-Wallis and post hoc Dunn's multiple comparisons test, further adjusted by the Benjamini-Hochberg FDR method.

(D) An emerging GC-like cell in a 2-dpf Q32 mYFP NICD Atoh7 retina with ptf1a morpholino (MO). The cell migrates basally, loses its bipolar morphology, and grows out an axon (arrowhead) and apically directed processes.

(E) Two Q32 GC-like cells in a 3-dpf transgenic embryo (Q32; mYFP; NICD; Atoh7 and ptf1a MO). Putative axons exit the retina (arrowhead). Gamma adjusted. (F) Dorsal view of a 3-dpf embryo (rostral top, eyes lateral) in which GC-like cells were induced (Q32; mYFP; NICD; Atoh7 and ptf1a MO), showing axons at the chiasm (arrowhead). Gamma adjusted.

Scale bars, 20 µm (A); 10 µm (D and E); 100 µm (F). GCL, ganglion cell layer; INL, inner nuclear layer; OLM, outer limiting membrane. See also Figure S4.



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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cub.2021.08.049.

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AUTHOR CONTRIBUTIONS

P.E., E.P., P.R.W., T.M., and L.G. conceived of the project and designed the experiments. L.G. and T.M. supervised the project. P.E. and E.P. performed the experiments. P.E., S.C.S., T.Y., and L.G. generated new constructs and transgenic lines. R.P. advised on the data analysis. L.G. wrote the paper, with input from all of the authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Mouse monoclonal anti-c-myc, clone 9E10, 1:100	Sigma-Aldrich	Cat# M5546; RRID: AB_260581
Chicken polyclonal anti-GFP, 1:1000	Abcam	Cat# ab13970; RRID: AB_300798
Mouse monoclonal anti-glutamine synthetase, clone GS-6, 1:50	Sigma-Aldrich	Cat# MAB302; RRID: AB_2110656
Goat anti-mouse Alexa 647, IgG1, 1:250	Invitrogen	Cat# A-21240; RRID: AB_141658
Goat anti-chicken Alexa 488, IgG (H+L), 1:250	Invitrogen	Cat# A-11039; RRID: AB_142924
Goat anti-mouse Alexa 568, IgG2a, 1:250	Invitrogen	Cat# A-21134; RRID: AB_2535773
Chemicals, peptides, and recombinant proteins		
N-[N-(3,5-Difluorophenacetyl)-L-alanyl]-S- ohenylglycine t-butyl ester (DAPT)	Enzo Life Science	Cat# 50-200-8542
Critical commercial assays	<u> </u>	
MyTaq Extract-PCR kit	Bioline	Cat# BIO-21126
Omniscript RT kit	QIAGEN	Cat# 205111
Ambion mMessage mMachine kit	Thermo Fisher Scientific	Cat# AM1340
Experimental models: Organisms/strains		
Zebrafish: Tg(vsx1:GFP)nns5: nns5Tg	12,16	ZFIN: ZDB-ALT-090116-1
Zebrafish: Tg(vsx2:GFP)nns1: nns1Tg	12,60	ZFIN: ZDB-ALT-061204-2
Zebrafish: Tg(UAS:gap43-YFP)q16b: q16Tg	61	ZFIN: ZDB-ALT-071129-3
Zebrafish: Tg(crx:MA-CFP)q20: q20Tg	21	ZFIN: ZDB-ALT-131118-1
ebrafish: Tg(vsx1:Gal4)q32: q32Tg	14	ZFIN: ZDB-ALT-170831-2
ebrafish: Tg(14xUAS:memTagRFP-T)	14	ZFIN: ZDB-ALT-170831-1
Zebrafish: Tg(ptf1a:eGFP)jh1: jh1Tg	19	ZFIN: ZDB-ALT-070531-2
Zebrafish: Tg(UAS:myc-notch-intra): kca3Tg	33	ZFIN: ZDB-ALT-020918-8
Zebrafish: Tg(T2KTp1bglob:hmgb1mCherry) h11: jh11Tg	28	ZFIN: ZDB-ALT-101006-1
Zebrafish: Tg(gfap:GFP)mi2001: mi2001Tg	34	ZFIN: ZDB-ALT-060623-4
Zebrafish: Tg(UAS:Kaede)s1999t: s1999tTg	62	ZFIN: ZDB-ALT-070314-1
Zebrafish: Tg(-17.6isl2b:GFP): zc7Tg	52	ZFIN: ZDB-ALT-100322-2
Zebrafish: Tg(-1.8ctbp2:gap43-EGFP) mb1: lmb1Tg	20	ZFIN: ZDB-ALT-120320-3
Zebrafish: Tg(EPV.TP1-Mmu.Hbb: nist2h2I-mCherry)s939: s939Tg	27	ZFIN: ZDB-ALT-110503-3
Zebrafish: Tg(14xUAS:mCherry)s1984t: s1984tTg	63	ZFIN: ZDB-ALT-130702-1
ebrafish: Tg(5xUAS:Atoh7)	This paper	N/A
Zebrafish: Tg(5xUAS:Ptf1a)	This paper	N/A
Dligonucleotides		
For all primers used in cloning, see Table S1	Metabion	N/A
For all primers used in genotyping, see Table S2	Metabion	N/A
Morpholino p53: 5'- GCGCCATT GCTTTGCAAGAATTG-3'	Gene tools, ⁶⁴	ZFIN: ZDB-MRPHLNO-070126-7
Morpholino:MO1-ptf1a: 5'-CCAACA CAGTGTCCATTTTTGTGC-3'	Gene tools, ^{43,46}	ZFIN: ZDB-MRPHLNO-070531-6
Recombinant DNA		
oCH-5xUAS:Ptf1a	This paper	N/A
oCH-5xUAS:Atoh7	This paper	N/A

(Continued on next page)





Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
atoh7 cDNA, I.M.A.G.E. clone	Source Bioscience	Cat# IRBVp5006D093D
pCS2 Numb-GFP	65	N/A
Software and algorithms		
ImageJ/Fiji	66	RRID: SCR_002285; https://fiji.sc/
Imaris	Bitplane	RRID: SCR_007370; https://imaris.oxinst.com/
Adobe Illustrator	Adobe	RRID: SCR_010279; https://www.adobe.com/
Adobe Photoshop	Adobe	RRID: SCR_014199; https://www.adobe.com/
GraphPad Prism	Graphpad Software	RRID: SCR_002798; https://www.graphpad.com

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Leanne Godinho (leanne.godinho@tum.de).

Materials Availability

Plasmids are available upon request to the lead contact.

Data and Code Availability

- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Experiments were performed according to local regulations (Regierung von Oberbayern). Zebrafish were maintained and bred as previously described. Embryos were kept in 0.3x Danieau's solution at 28.5°C and staged as previously described. Eish were either in an AB wild-type, Tuebingen Long Fin (TLN) or roy orbison background. All experiments were performed on animals between 2 and 5 days post fertilization (dpf). During this period zebrafish are not sexually differentiated. The transgenic lines used are listed in the key resources table. Note that line Tg(vsx1:Gal4)q32 (Q32) was referred to as Tg(vsx1:Gal4)q26 (Q26). To generate the line, a 3.2 kb fragment upstream of the vsx1 coding sequence was used to drive the expression of Gal4-VP16. Crossing the Q32:Gal4 line, in conjunction with a uas:memTag-RFP-T line, to vsx1:GFP revealed that the Q32 driver faithfully labels a subset of vsx1:GFP+ cells. We generated Tg(5xuas:Ptf1a) and Tg(5xuas:Atoh7) by Tol2 mediated insertion.

METHOD DETAILS

Generation of constructs

To generate *pCH-5xuas*:Atoh7, the coding sequence of zebrafish *atoh7* (*atonal bHLH transcription factor 7*) was PCR amplified using primers containing EcoRl and NotI restriction sites (sequences of primers Atoh7 Forward and Atoh7 Reverse are available in Table S1) from a plasmid containing full-length *atoh7* cDNA (I.M.A.G.E Clone IRBVp5006D093D, Source Bioscience). The amplified *atoh7* sequence was cloned into a *5xuas* backbone using EcoRl and NotI. The *5xuas* backbone was generated by excising TagRFP-T out of *5xuas*:TagRFP-T (gift from Dr. M. Meyer, King's College, London, UK) using EcoRl and NotI. Subsequently, *5xuas*:Atoh7 was excised with Asel and AfIII and ligated into the pColdHeart Tol2 vector (gift from Dr. M. Nonet, Washington University, St. Louis, USA) using blunted Nhel and Clal restriction sites.

To generate *pCH-5xuas*:Ptf1a, the coding sequence of zebrafish *ptf1a* was amplified using primers containing Xmal and Notl restriction sites (sequences of primers Ptf1a Forward and Pf1a Reverse in Table S1) from zebrafish cDNA (obtained by reverse transcription, Omniscript RT kit, QIAGEN) and cloned into the *5xuas* backbone using Xmal/Notl. *5xuas*:Ptf1a was released with Asel and AfIII and ligated into the pColdHeart Tol2 vector using blunted Nhel and Clal restriction sites.

Capped RNA synthesis and injections

The PCS2 Numb-GFP plasmid was linearized using Notl. Capped mRNA was synthesized using the Ambion mMessage mMachine kit (Applied Biosystems) according to the manufacturer's instructions. Numb RNA was injected in one- to two-cell stage fertilized eggs at a concentration of 20ng/μl.





Mounting zebrafish larvae for in vivo imaging

Embryos were prepared for imaging as described previously. 14,72 Between 10 and 24 h post-fertilization (hpf), embryos were transferred to 0.3 × Danieau's solution containing 0.003% 1-phenyl-2-thiourea (PTU, Sigma) to inhibit melanin formation. Embryos were manually dechorionated (when necessary), 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) in glass covered 35mm dishes (MatTek, P35G-0-14-C).

In vivo imaging

Fish were imaged on an Olympus FV1000 confocal/2-photon, Olympus FVMPE-RS 2-photon, Olympus FV3000 or a Leica TCS SP8 microscope using water-immersion objectives (Olympus 20x/NA 0.95, Olympus 25x/NA 1.05, Zeiss 40x/NA 1.0, Nikon 25x/NA 1.1 and Nikon 40x/NA 0.8, Leica 25x/NA 0.95) or a silicon-immersion objective (Olympus 30x/NA 1.05). Embryos were maintained at 28.5°C during in vivo time-lapse recordings.

Photoconversion

The 'tornado scan' function on an Olympus FV1000 confocal was used to photoconvert Kaede in a local patch of Q32 cells within the INL of Q32; uas: Kaede double transgenic fish with a 405 nm laser. Retinae were imaged immediately following photoconversion (Day 0). Fish were then unmounted from agarose and maintained in the dark at 28.5°C until subsequent imaging time-points at 24h (Day 1) and 48h (Day 2).

Morpholino injections

An antisense ptf1a translation-blocking morpholino (0.5mM, MO1, Gene tools, for sequence see key resources table), was injected using a picospritzer into the yolk of one or two-cell stage compound transgenic embryos (Q32; mYFP; NICD; Atoh7). A p53 morpholino (0.02 to 1.0 mM, Gene tools, for sequence see key resources table) was injected into embryos from the vsx1:GFP line that were subsequently treated at 2 dpf with DAPT (see below).

DAPT treatment

N-[N-(3,5-Difluorophenacetyl)-L-alanyl]-S-phenylglycine t-butyl ester (DAPT) was used at a final concentration of 50 μM in 0.3x Danieau's containing 1% DMSO. Embryos were injected with a p53 morpholino to ameliorate DAPT induced toxicit. 64 At 2 dpf, embryos were transferred into DAPT containing medium (or DMSO containing medium as a control) and incubated for approximately 20 h before analysis.

Immunostaining

Immunostaining to detect glutamine synthetase, c-myc and YFP was performed on whole-mount compound transgenic embryos. Zebrafish embryos were fixed in 4% (wt/vol) paraformaldehyde (PFA) in 1x PBS at 4°C overnight. Following fixation, embryos were washed (3x5min) in 1x PBS, pH 7.4, and then incubated in 0.25% Trypsin in 1x PBS on ice for 3-4 min. Embryos were then washed 3-4 times in 1x PBS and incubated in 0.4% Blocking Reagent (Roche) in PBS for 1.5h. Incubation in primary antibodies, diluted in 1x PBS containing 1% Triton (PBST) and 0.4% Blocking reagent, proceeded for two days at 4°C. Following several washes in PBST for 4-5h, embryos were incubated in appropriate Alexa dye-coupled secondary antibodies overnight at 4°C in 0.4% Blocking reagent in PBST. Following several washes in 1x PBS, embryos were mounted in low-melting agarose (0.7%-0.8%, Sigma) and imaged. All incubation and washing steps were done on a rotating shaker.

Genotyping of Q32 crosses

The presence of the uas:NICD transgene was initially determined by immunostaining for the c-myc tag (99.1 ± 0.6% of YFP+ cells were c-myc+; 428 cells, 3 Q32 NICD fish). The presence of c-myc staining correlated perfectly with exuberant neuritic arbors and the loss of lamination in the IPL as well as supernumerary ACs in retinae from putative Q32 NICD fish (12 out of 12 fish). Thus, the presence of the NICD transgene could be readily determined by retinal morphology. The presence of the uas:Ptf1a or uas:Atoh7 transgene in embryos could be confirmed by cyan fluorescence in the heart as both constructs were generated using the pCold-Heart Tol2 vector in which cmcl2 promoter elements drive the expression of CFP.

For some experiments we genotyped individual embryos after confocal imaging to definitively identify carriers of the uas:NICD or uas:Atoh7 transgenes. Between 6-11dpf, fish were anesthetized in 0.2% tricaine (PharmaQ) and processed for genotyping. DNA extraction was performed using a MyTaq Extract-PCR kit (Bioline, BIO-21126) according to the Manufacturer's guidelines. In brief, each reaction required 20μl Buffer A, 10μl Buffer B and 70μl Nuclease-Free Water (Ambion). Each larva was placed into a clean 1.5ml tube and the extraction mix was added. The reaction was incubated for 75°C for 5-6min and vortexed at least twice. Enzymatic deactivation was achieved by heating to 95°C for 10min followed by centrifugation at 14000rpm for 5min. The genomic DNA of each individual larva was transferred to a clean 1.5ml tube and used as a template for the genotyping. PCR to detect the transgenes used OneTag hot Start DNA Polymerase (NEB, M0481) along with primers to detect Actin as a proxy to determine the integrity of the genomic DNA. Primers used for genotyping are listed in Table S2.



QUANTIFICATION AND STATISTICAL ANALYSIS

Cell type classification of Q32 cells

Q32 cells were classified as BC or AC based on neurite morphology (presence or absence respectively of an apical process) and position of the soma (lower or upper half of the INL). To obtain the AC-BC ratio of different Q32 genotypes, a maximum intensity projection (MIP) of a few imaging planes (typically 5) was obtained, and all Q32 cells (at least 70 per retina) in the MIP were classified. Q32 divisions were classified as BC-BC, AC-BC or AC-AC based on the criteria described above.

Quantification of Kaede recovery in photoconverted Q32 cells

The recovery of green fluorescence intensity of Kaede protein (I_{CV}) was measured in a region of interest (ROI) encompassing the soma of individual BCs and ACs in a single image plane of a confocal stack and the background fluorescence (I_{BG}) averaged from six regions (Q32 negative) was subtracted to correct for auto-fluorescence and potential out-of-focus contributions. In order to compare across different samples, fluorescence intensity values were normalized to the average fluorescence intensity of 6 unconverted BCs (I_{UC}) from the same section. The recovery of green fluorescent Kaede protein in Q32 BCs and ACs was expressed as a percentage:

Recovery of green fluorescence in Q32 BC =
$$\frac{I_{CV_BC} - I_{BG}}{I_{UC_BCs} - I_{BG}}\%$$

Recovery of green fluorescence in Q32 AC =
$$\frac{I_{\text{CV_AC}} - I_{\text{BG}}}{I_{\text{UC_BCs}} - I_{\text{BG}}}\%$$

The unconverted BCs were chosen in the vicinity of the photoconverted area and ± 5 sections from the depth in which the photoconverted area was in focus. The same procedure was followed to obtain background values.

Analysis of DAPT treated animals

The effect of DAPT treatment was quantified in two independent quadrants of the peripheral retina of vsx1:GFP fish. In a given quadrant, the total number of *vsx1*:GFP⁺ ACs were counted in 10 z-planes.

Notch-Reporter levels in differentiated Q32 cells

Tp1:hmgb1mCherry fluorescence intensity levels were determined in Q32 ACs and their potential sibling BCs. If at least one of the Q32 cells had a fluorescence intensity level twice above background, the cell group was used for analysis (34/77 cell groups could be used). To classify a cell group as 'AC high', 'BC high' or 'equal', Notch-Reporter levels in the Q32 AC were compared to the surrounding Q32 BCs with the highest Notch-Reporter fluorescence. If the Q32 AC had twice the brightness of the Q32 BC, the group was classified as 'AC high'. Similarly, in cases in which the Q32 BC had twice the brightness of the Q32 AC, the group was classified as 'BC high'. When no cell had at least twice the fluorescence intensity of the other, the group was classified as 'equal'.

Monitoring Notch-Reporter levels in time-lapse recordings of the Q32 lineage

Tp1:H2BmCherry fluorescence intensity levels were monitored in the Q32 lineage by in vivo imaging. Q32 BC-AC pairs were identified and their somata were manually demarcated based on their mYFP expression in a single image plane of a confocal stack using the ImageJ freehand tool. Notch-Reporter levels were measured based on the fluorescence intensity of the mCherry channel within the demarcated areas. The same approach was used for Q32 transdifferentiation events in the NICD OE condition. Background values (mCherry channel) from three areas at the same time point were averaged and subtracted from the fluorescence intensity values of the cells being measured. Contributions of fluorescence from cells outside of the Q32 lineage were identified as moving "objects" and used to correct the demarcation of each cell of interest accordingly. Finally, the fluorescence values were normalized to the time point before mitosis for the Q32 BC-AC siblings or to the first time point seen for the transdifferentiation events after background subtraction.

Division rate analysis

The number of mitotic divisions in the Q32 lineage in WT and NICD OE, irrespective of their outcome (BC-AC or BC-BC), were counted over a time span of ~18h between 52 hpf and 70 hpf. The division rate was expressed as the total number of divisions per hour.

Cleavage plane analysis

The cleavage planes of dividing progenitors that ultimately generated BC-AC daughters were analyzed. Cell divisions were analyzed in imaging volumes with a reference to the apical surface. Divisions were classified as apico-basal if the cells divided perpendicular to the apical surface and both siblings were present in the same imaging plane. Circumferential divisions occurred parallel to the apical surface and when both siblings were present in the same imaging plane. Central-peripheral divisions also occurred parallel to the



apical surface but the two cells were at different z-depths (larger than 2.4 µm) representing the central-peripheral axis of the retina. When dividing cells could not be strictly categorized in these three categories we ascribed them to an intermediate class.

Statistics

The Shapiro-Wilk test was used to determine whether parametric or non-parametric tests were appropriate for determining statistical significance with GraphPad Prism 8. The specific statistical tests used to compare data-sets are indicated in the respective part of the text or figure legend. P \leq 0.05 denoted with '*', \leq 0.01 with '**', \leq 0.001 with '***', \leq 0.0001 ****.