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Biasing amacrine subtypes in the Atoh7-lineage through expression of Barhl2

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Abstract

Within the developing vertebrate retina, particular subtypes of amacrine cells (ACs) tend to arise from progenitors expressing the bHLH transcription factor, Atoh7, which is necessary for the early generation of retinal ganglion cells (RGCs). All ACs require the post-mitotic expression of the bHLH transcription factor Ptf1a, however Ptf1a alone is not sufficient to give subtype identities. Here we use functional and *in vivo* time-lapse studies in the zebrafish retina to investigate on the developmental programs leading to ACs specification within the subsequent divisions of Atoh7-positive progenitors. We find evidences that the homeobox transcription factor Barhl2 is an AC subtype identity-biasing factor that turns on within Atoh7-positive descendants. *In vivo* lineage tracing reveals that particular modes of cell division tend to generate Barhl2-positive precursors from sisters of RGCs. Additionally, Atoh7 indirectly impacts these division modes to regulate the right number of *barhl2*-expressing cells. We finally find that Atoh7 itself influences the subtypes of Barhl2-dependent ACs. Taken together, our study uncovers lineage-related and molecular logic of subtype specification in the vertebrate retina, by showing that specific AC subtypes arise *via* a particular mode of cell division and a transcriptional network cascade involving the sequential expression of first *atoh7* followed by *ptf1a* and then *barhl2*.

Keywords

fate determination; cell lineage; Barhl2; subtype specification; retina; zebrafish

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Authors' contributions LP conceived this study, supervised the project and wrote the manuscript. PRJ and SA contributed equally to this work, designing and carrying out experiments and analyses. Experiments by PRJ were conducted in the laboratory of and supported by PC. PRJ carried out experiments shown in Figures 1, 2, 4, 6, 7 and Movie 1 and wrote the first manuscript draft with LP. Early-stage experiments by PRJ were conducted in the laboratory of WAH. All experiments by SA, AP and LP were carried on in the laboratory of LP. SA carried out the cloning and experiments shown in Figures 3 - 5, 7 - 9 and Movie 3. SA also performed caspase cell death experiments. AP and LP carried out lineage experiments and cell tracking (Movie 2, 4 and Figure 7). FA and SH generated the transgenic lines used for this study. All authors revised and approved the final manuscript.

INTRODUCTION

A major challenge in vertebrate neurobiology is to understand how the developmental programs of a neural progenitor cell are regulated *in vivo* in the context of cell lineages and modes of cell division. Within the vertebrate retina, some types of neurons tend to be lineally related or descendants of common progenitor cells (Poggi et al., 2005b; Vitorino et al., 2009; Feng et al., 2010; Brzezinski et al., 2011; Jusuf et al., 2011). The bHLH transcription factor *Atoh7* (a.k.a *Ath5*) is required for RGCs development (Brown et al., 2001; Kay et al., 2001; Vetter and Brown, 2001; Wang et al., 2001; Ghiasvand et al., 2011), and turns on just before mitosis that precedes their birth (Poggi et al., 2005b). One cell from this mitosis differentiates as a RGC. However, many other cell types, including some subtypes of ACs also come from *atoh7* expressing progenitors (Poggi et al., 2005b; Feng et al., 2010; Jusuf et al., 2011). The sisters of RGCs must therefore generate these other cell types.

The fates of all retinal neurons that primarily express the inhibitory neurotransmitters GABA or glycine (horizontal cells and ACs) require the expression of the Pancreas transcription factor 1a (*Ptf1a*) (Fujitani et al., 2006; Dullin et al., 2007; Nakhai et al., 2007; Jusuf et al., 2011). All ACs express *Ptf1a*, but *Ptf1a* alone is not sufficient to confer subtype-specificity (Jusuf et al., 2011). However, precursors that express both *atoh7* and *ptf1a* tend to differentiate into specific subtypes of ACs, thus suggesting that other key factors might regulate AC subtypes within this lineage (Jusuf et al., 2011).

Barhl homeobox transcription factors have been implicated in ACs diversity and RGC development downstream of *Atoh7* (Poggi et al., 2004; Ding et al., 2009). Targeted disruption of *barhl2* alters AC subtype composition and survival of RGCs (Ding et al., 2009). Nothing is known on the lineage-origin of *barhl2*-expressing cells, the networks in which *Barhl2* specifies AC subtypes, or how it works in relation to genes that drive the same (*ptf1a*) or alternate fates (*atoh7*). In zebrafish, additional whole genome duplication has generated another *barhl* paralog (Reig et al., 2007; Schuhmacher et al., 2011). *Barhl1.2* is specifically expressed in RGCs, while *barhl2* is expressed in ACs (Schuhmacher et al., 2011). This led us to investigate the distinct role of *Barhl2* as an AC subtype-biasing factor downstream of *Atoh7*. We found that *barhl2*-expressing precursors arise within the *Atoh7*-lineage. *Barhl2* expression, however, does not depend on *Atoh7*, but on *Ptf1a*, and is necessary and sufficient for biasing AC subtypes. Additionally, *Atoh7* affects the identities of *Barhl2*-dependent ACs. With timelapse imaging (Poggi et al., 2005b; Poggi et al., 2005a) we traced the origins of *Barhl2*-positive cells. We found that these cells arise as one of the two post-mitotic daughters of a dividing RGC's sister, i.e. *Barhl2* ACs tend to be nieces of RGCs. Our study provides *in vivo* evidences that modes of cell division and lineage-restricted cell fate determination programs regulate the correct number of neuronal subtypes within particular progenitor pools.

MATERIALS AND METHODS

Animals and ethics statements

Zebrafish breeding / raising followed standard protocols. Fish were maintained at 26.5°C and embryos raised at 28.5°C or 32°C and staged as described (Kimmel et al., 1995). Fish were housed in three facilities: Fish facility of our German laboratory (built in accordance to Tierschutzgesetz 111, Abs. 1, Nr. 1 and with European Union animal welfare guidelines); fish facility at the University of Cambridge, UK; and FishCore at Monash University, Australia. Each facility is under supervision of and in accordance with local animal welfare agencies. Zebrafish (*Danio rerio*) embryos of either sex were used exclusively prior to free-

feeding stages. Embryos used for whole-mount imaging were treated with 0.0045% 1-Phenyl-2-Thiourea (PTU) (Sigma, Gillingham, Dorset, UK) to delay pigment formation.

Fish lines

Seven transgenic lines expressing *GFP*, *dsRed*, *gap43-GFP* or *gap43-RFP* under the control of different promoters were used in this study: Tg(*barhl2:GFP*) line (Kinkhabwala et al., 2011); Tg(*ptf1a:GFP*) line (Godinho et al., 2005) kindly provided by Steven D. Leach (John Hopkins Medical Institutions, Baltimore, USA); Tg(*atoh7:gap43-GFP*), Tg(*atoh7:GFP*), Tg(*atoh7:gap43-RFP*) and Tg(*atoh7:gal4/pUAS:gap43-GFP*) lines (Zolessi et al., 2006). For the Tg(*ptf1a:dsRed*) line (Tg(-5.5*ptf1a:DsRed*)_{ia6}) we created a plasmid containing 5.5 kb of the 5' region of the *ptf1a* gene cloned upstream of DsRed2 in pT2AL200R150G vector (Kawakami, 2004). The plasmid was injected with Tol2 transposase mRNA and F1 progeny of different insertion lines was screened. The *ia6* allele faithfully represents the endogenous expression of *Ptf1a* mRNA and has a comparable expression pattern to that of the previously characterised Tg(*ptf1a:GFP*) with the *dsRed* showing only a slight delay in expression (data not shown). Double transgenic lines were generated via outcrossing.

Morpholino injection

Translation blocking morpholino oligonucleotides (MO) obtained from Gene Tools, LLC (Oregon, USA) were reconstituted as 1mM stock solutions in water and injected into the yolk of 1 - 2 cell stage embryos. A MO targeted against a region 44 bp upstream from the translational start site with sequence 5'-TTGCCAGTAACAACAATCGCCTAC-3' was used to knockdown *ptf1a* (10 – 12ng / embryo) (Lin et al., 2004; Jusuf et al., 2011). A MO with sequence 5'-TTCATGGCTCTTCAAAAAGTCTCC-3' was used to knockdown *atoh7* (Pittman et al., 2008). A *barhl2* translation MO targeting 6 bp upstream from the translational start site with the sequence 5'-AGAAAAGGATGAGCACTCAAGTCGT-3' was designed and injected at 0.5mM / embryo. Injections of standard control morpholino with sequence 5'-CCTCTTACCTCAGTTACAATTTATA-3' up to 12 ng had no effect. The 5 bp *Barhl2* mismatch MO with sequence 5'-AGAATACGATCAGCACTGAACTCGT-3' is also comparable to uninjected (**data not shown**). Cell-autonomous role of *Barhl2* was assessed using transplantation technique. Briefly, 10 – 20 cells were transplanted from blastula stage donors (cells labelled with *H2A-GFP* or *H2B-RFP*) into the animal poles of blastula stage host embryos. Integration and survival of transplanted cells was aided by injecting p53 MO with sequence 5'-GCGCCATTGCTTTGCAAGAATTG-3' into donor embryos. Retinas injected with standard control MO together with p53 MO display normal expression of *Barhl2* protein as shown by antibody staining (**data not shown**).

Overexpression plasmid cloning

The coding sequence of *barhl2* was PCR amplified using 5'-ATGGAAGGATCCAGTGGGGCTAGT-3' and 5'-CCGAGCATGCGGTGTGCC-3' for which the reverse primer was tagged with the T2A sequence 5'-AGGGCCGGGATTCTCCTCCACGTCACCGCATGTTAGAAGACTTCTCTGCCCTC-3' (Kim et al., 2011). *H2B-RFP* fused coding sequences were PCR amplified using 5'-ATGCCAGAGCCAGCGAAGTCT-3' and 5'-GATGTACACGGCGCCGGT-3' primers for which the forward primer was tagged with the T2A sequence 5'-GAGGGCAGAGGAAGTCTTCTAACATGCGGTGACGTGGAGGAGAATCCCGGCC-3'. A fusion PCR was performed to generate the *barhl2-t2a-h2b-rfp* product, which was cloned into a pUAS vector containing 16 cassettes of the Upstream Activation Sequence and recognition sequences for I-SceI meganuclease for efficient transgenesis.

Immunohistochemistry

Primary antibodies were diluted in blocking solution: rabbit anti-calretinin (Chemicon AB5054; 1:1000), rabbit anti-Sox2 (Chemicon AB5603, 1:200), mouse anti-parvalbumin (Chemicon MAB1572, 1:1000), rabbit anti-GABA (Sigma A2052, 1:500), rabbit anti-calbindin (Calbiochem PC253L, 1:500), rabbit anti-serotonin (Sigma S5545, 1:50), mouse anti-tyrosine hydroxylase (Millipore MAB 318, 1:100), rabbit anti-Neuropeptide Y (Immunostar 22940, 1:500), rabbit anti-Barhl2 (Santa Cruz Biotechnology sc-68370, 1:50 for immunohistochemistry, 1:1000 for Western blot). Secondary antibodies were goat or donkey anti-mouse, anti-rabbit or anti-goat IgG conjugated to Alexa 488, 546, 594 or 647 fluorophores (1:1,000 - 1:2,000; Molecular Probes, Eugene, OR, USA).

For most antibodies, embryos were fixed in 4% paraformaldehyde (PFA) in 0.1M PB overnight (maximum 2 hours for ChAT immunohistochemistry) at 4°C, rinsed, cryoprotected in 30% sucrose, embedded in OCT and cryosectioned at 14µm thickness. For GABA immunohistochemistry, embryos were fixed in 4% PFA / 0.05% glutaraldehyde, 5mM EGTA, 5mM MgSO₄, 0.1% triton-X100 in 0.1M PB for 3 hours at room temperature. All staining steps are performed at room temperature unless stated otherwise. For Sox2 immunohistochemistry antigen retrieval was performed by immersing sections in 0.01M sodium citrate buffer, pH 6.0 at 95 °C for 10 min prior to blocking. All sections were incubated in blocking solution (10% heat-inactivated goat serum (HIGS), 1% bovine serum albumin, 0.2% Triton X-100 in PBS) for 30 minutes (sections) or 60 minutes (wholamounts). For staining with goat anti-ChAT antibody, sections were blocked in 10% donkey serum instead. Sections were incubated in primary antibodies overnight, secondary antibodies for 60 minutes, and nuclei were counterstained with 4',6-diamidino-2-phenylindole (DAPI). Sections were mounted with Fluorosave (Calbiochem, Merck Chemicals Ltd., Nottingham, UK) or Mowiol.

Whole-mount single and fluorescent *in situ* hybridization

In situ mRNA hybridization was performed as described (Lin et al., 2004). The full-length cDNA *barhl2* sequence was subcloned from a zebrafish IMAGE clone (IMAGE: 7452725; IRBOP991F0870D, Source BioScience UK Limited, Nottingham, UK) in pME18S-FL3 into a pCS2+ vector to generate digoxigenin- and fluorescein-labelled riboprobes. For antisense probe, we linearized with NotI (Fermentas or New England Biology) and transcribed with Sp6 (mMessage mMachine® Sp6, Ambion). For sense probe, we linearized with BamHI (Fermentas or New England Biology) and transcribed with T7 (mMessage mMachine® T7, Ambion). *Atoh7* probes were generated as described (Schuhmacher et al., 2011). *Ptf1a* probes were generated directly by RTPCR (one step RTPCR kit, Qiagen) using total mRNA extracted from zebrafish embryos of 50 hpf and 5' TTCGAGAGACCACTTGGACA3' forward primer and T7 tailed 5' - CCAAGCTTCTAATACGACTCACTATAGGGAGAGGCTGAAACACAGATAGTCACA A-3' reverse primer. Single probe *in situ* hybridization was done as described (Thisse and Thisse, 2008) with minor modifications. Embryos underwent a stepwise dehydration series into 100% methanol and subsequent rehydration into 0.1% Tween in PBS. Permeabilization was achieved using age-dependent concentrations of proteinase K treatment at room temperature, followed by postfixation in 4% PFA in PBS. After prehybridization, hybridization with digoxigenin-UTP labelled probes (Roche Applied Science) was performed overnight at 65°/68 °C. Signal was detected with nitroblue tetrazolium chloride/ 5-bromo-4-chloro-3-indolyl-phosphate, toluidine salt (NBT/BCIP BM Purple, Roche Products Ltd., Welwyn Garden City, UK).

For double fluorescent whole mount *in situ* hybridization (FISH) standard digoxigenin- and fluorescein-labelled riboprobes were combined with Tyramide Signal Amplification (TSA),

as described in Schumacher et al., 2011. Riboprobes were incubated for 30 minutes (*barhl2*), 40 minutes (*atoh7*) or 24 hours (*ptf1a*). Embryos were kept in the dark for following steps. Embryos were washed with TNT (0.1M Tris pH7.5, 0.15M NaCl, 0.1% Tween20), incubated with 1% H₂O₂ in TNT for 20 minutes, washed several times and blocked with TNB (2% DIG Block [Roche] in TNT) for 1 hour followed by incubation with anti-digoxigenin-POD Fab fragments (Roche, 1:50 in TNB). Signal was detected using Fluorescein-Tyramide (FITC), Cyanine3-Tyramide (Cy3) or Cyanine5-Tyramide (Cy5) fluorophores (PerkinElmer). Embryos were incubated in DAPI in TNT overnight at 4°C and washed with TNT.

Imaging

NBT/BCIP stained embryos were mounted in 87% glycerol and imaged with Leica DM5000B compound microscope at 10x or 20x. Images were acquired with a Leica CD500 camera using Leica FireCam 1.7.1. FISH embryos were mounted on 100 × 15 mm glass bottom Petri dishes using 1% low melting agarose and z-stacks taken at the Leica SP5 confocal microscope.

Images of fixed and *live* embryos were acquired on a dissecting stereomicroscope equipped with epi-fluorescence (Leica MZ FLIII). Photomicrography of whole-mount eyes or sections was performed with a laser confocal system (Leica TCS-NT, Leica SpE or Leica Sp5 confocal laser scanning microscopes using a Leica 40x, 1.2 NA or Leica 63x, 1.2 NA water immersion objectives) or Nikon fluorescence microscopes, equipped with cooled charge-coupled device (CCD) Hamamatsu Orca cameras and automated z-drive and fluorescence shutters.

At the confocal, excitation was achieved with following laser lines: 405nm (DAPI), 488nm argon (GFP, Alexa 488), 568nm (RFP, DsRed, Alexa 546), 594nm (Alexa 594) and 633nm (Alexa 647). Images were taken through whole-mount fixed and *live* embryos as described (Poggi et al., 2005b). Sequential image acquisition was performed with emission detected at 500-550nm (FITC), 650-700nm (Cy3), 650-800nm (Cy5) and 400-500nm (DAPI) using individual descanned PMT detectors. Optical sections (40 – 60µm for timelapse and < 100µm for fixed embryos) of 1 µm thickness were taken and Kalmann averaged 2 or 4 times. For time-lapse, images were taken every 5 or 10 minutes for 24 – 42 hours. Motorized XY stage was used to image multiple embryos. Laser power was minimized to avoid bleaching and phototoxicity.

Image processing and cell tracking

Image data was acquired using Leica Application Suite (LAS), Leica TCS NT or Leica LCS software, processed and analysed using Volocity Analysis version 5.3 (Improvision, Coventry, UK). Brightness and contrast were adjusted with Adobe Photoshop CS3 and CS4. Cell tracking was performed using the Volocity classification module (Improvision). Double-labelled *Barhl2:GFP/Atoh7:gap43-RFP* cells were randomly selected and *Atoh7:gap43-RFP* positive cells tracked backward in time.

Analysis

Numbers used for each analysis and age (generally 4 or 5 dpf) are indicated in the result section or figure legend. The majority of quantification was performed in the central retina in which the relatively older neurons express the proteins recognised by the antibodies. The region was defined by drawing a straight line through the centre of the lens just central to the ciliary margin as described previously (Holt et al., 1988). The half covering the retina was subdivided into four segments and quantification performed in the central two. For transplanted cells of rare ACs, the whole retina was included. For markers with < 10 cells

per section, 5 sections were combined (i.e. serotonin, tyrosine hydroxylase and neuropeptide Y). Subtypes of *barhl2*-expressing AC subtypes were classified morphologically as described previously (Jusuf and Harris 2009).

Statistical analysis

Statistical tests were performed using Prism software with $p < 0.05$ used as criterion level. Comparison of morphologically characterised subtypes from *barhl2*-expressing cells: Fisher's exact test. Tests used for *atoh7* knock down experiment: unpaired t-test with Welch's correction, Gaussian distribution, 2-tailed (GFP and GABA quantification); Fisher's exact test, 2-sided and $\alpha < 0.05$ (Serotonin and NY quantification). Tests used for *barhl2* over-expression experiment with GABA: Mann-Whitney test, Gaussian approximation, 2-tailed. For *barhl2* knockdown transplantation experiments, the binomial test was used, as the number of cells per image was too low to compare means.

RESULTS

Zebrafish *barhl2* is expressed differentially in subtypes of inhibitory cells

To investigate the possibility that the Barhl2 paralog in the zebrafish plays a specific role as an amacrine subtype specification factor within the Atoh7-lineage, we first assessed its gene expression with respect to the *pancreas transcription factor 1a* (*ptf1a*). Ptf1a specifies all inhibitory (defined here as primarily expressing the inhibitory GABA or glycine neurotransmitters) neurons (ACs and HCs) and is excluded from excitatory (defined here as expressing the excitatory glutamate neurotransmitter) cells (Jusuf and Harris, 2009). We used the double transgenic line Tg(*ptf1a:dsRed/barhl2:GFP*) to assess whether cells expressing *barhl2* also express *ptf1a*. Quantification in 4 days post-fertilization (dpf) Tg(*ptf1a:dsRed/barhl2:GFP*) transgenic embryos showed that Barhl2:GFP cells co-labelled with Ptf1a:DsRed signal ($94.9\% \pm 0.5\%$ SEM) primarily in ACs (Figure 1). Additionally, the few Barhl2:GFP cells within the ganglion cell layer (GCL) co-expressed Ptf1a:DsRed, suggesting that these are displaced ACs. One striking observation was that, although all Barhl2:GFP cells are Ptf1a-positive, only about $58.5\% (\pm 1.2\%$ SEM) of Ptf1a-expressing ACs are Barhl2:GFP-positive, suggesting that Barhl2 marks a subpopulation of Ptf1a-derived ACs (Figure 1). We found that some Ptf1a:DsRed-positive HCs are also Barhl2:GFP-positive (Mo et al., 2004; Ding et al., 2009). These cells however are mainly located in the retinal periphery, and in these HCs, Barhl2:GFP expression was highly variable (data not shown). Thus, stable *barhl2* paralog expression in the zebrafish retina is largely restricted to ACs.

As only 58.5% of Ptf1a-positive ACs turn on Barhl2:GFP, we wondered whether *barhl2* is expressed in specific subtypes. Previous studies implicated Barhl2 in biasing specific AC identities (Yazulla and Studholme, 2001; Clemente et al., 2004), but a detailed characterisation of subtypes expressing *barhl2* and how this correlates with changes during loss- or gain-of-function studies are still missing. We therefore first performed immunohistochemical staining with nine antibodies in 5 dpf Tg(*barhl2:GFP*). We specifically chose a range of different markers. Serotonin, tyrosine hydroxylase, neuropeptide Y, and choline acetyltransferase mark non-overlapping individual subtypes based on co-labelling or neurite morphology. Calbindin, parvalbumin and calretinin are calcium-binding markers that may label more than one subtype (e.g. parvalbumin labels 2 subtypes non-overlapping with calretinin labelled cells) (Yeo et al., 2009). Finally, we chose the more general marker GABA, which labels half of the zebrafish amacrine cells (Jusuf and Harris 2009) and likely overlap with some of the subtype specific markers (e.g. serotonin). We found that the vast majority of serotonin, calbindin and GABA-expressing AC types express Barhl2:GFP either strongly (serotonin) or at medium levels (calbindin, GABA;

Figure 2A – C, and J), in line with GABAergic subpopulations overlapping with serotonin and/or calbindin labelled subtypes. Calretinin, tyrosine hydroxylase or parvalbumin-expressing amacrine subtypes had more variability with roughly half of the labelled ACs expressing some Barhl2:GFP (Figure 2D – F, and J). Amacrine subtypes labelled by neuropeptide Y, Sox2 or choline acetyltransferase very rarely expressed Barhl2:GFP and never at high levels (Figure 2G – I, and J). These observations are consistent with the hypothesis that Barhl2 may be involved in the differentiation of specific AC subtypes in the zebrafish retina and that differences in expression levels may be important in biasing subtype fates.

Barhl2 functions downstream of Ptf1a to bias amacrine subtypes identities

As *barhl2* is restricted to some AC subtypes, its expression might be turned on downstream of Ptf1a only in certain AC precursor populations to specify their identities. To first temporally locate *barhl2* within the transcriptional cascade that leads to AC specification *in vivo*, we analysed the dynamics of its expression at the cellular level with respect to *ptf1a*. The first Barhl2:GFP signal starts at 35 hpf and it faithfully recapitulates the endogenous *barhl2* mRNA expression as revealed by double *in situ* hybridization with *GFP* mRNA (Figure 3A, B) or GFP protein staining (Figure 3C, D). 3D time-lapse imaging starting from 30-32 hours post-fertilization (hpf) show the first Barhl2:GFP signal at 35 hpf (Movie 1). These Barhl2:GFP positive cells were never seen dividing, always being either in the process of or having finished migrating basally to the inner nuclear layer where ACs reside (Movie 1). Whole mount double fluorescent *in situ* hybridization shows co-expression of *barhl2* and *ptf1a* mRNAs in individual cells (Figure 4A, B). Cells expressing *ptf1a* only can be observed apically, suggesting that as cells migrate basally to the future AC layer, they first express *ptf1a*. Consistently, our 3D *in vivo* time-lapse imaging shows that within individual developing neurons, Ptf1a:DsRed is turned on apically just after mitosis (Jusuf et al., 2011). In some of these cells Barhl2:GFP is turned on when they have reached the future AC layer (Figure 4C – C’). We tested directly, if Barhl2 functions downstream of Ptf1a, by injecting Ptf1a translational MO or control MO into Tg(*barhl2:GFP*) embryos. The Ptf1a morphants show a drastic loss in the number of Barhl2:GFP cells (Figure 4D, E) including ACs (59.1 ± 1.2 SEM to 9.02 ± 0.82 SEM; p – value < 0.0001) (Figure 4F). Because Ptf1a morphant cells remain within the retina and are re-specified as excitatory cells (Jusuf et al., 2011), these results demonstrate that *barhl2* turns on in inhibitory cell precursors downstream of Ptf1a.

We assessed how the loss of Barhl2 affects the development of *ptf1a*-expressing neurons. We used a morpholino (MO), which effectively knocked down Barhl2 protein translation as shown by western blot (50 hpf embryos) and antibody staining in hindbrain and retina (72 hpf embryos) (Figure 5). Standard control morpholino injected embryos were comparable to WT (Figure 5F’), as were standard control MO + p53 MO and 5 bp mismatch MO (data not shown). As recent studies implicated Barhl2 in cell survival (Ding et al., 2009; Juraver-Geslin et al., 2011), we firstly assessed activated Caspase-3 immunostaining in Barhl2 morphants, which revealed only a small non-significant increase in retinal apoptosis including in the AC layer (INL, p – value = 0.3; data not shown). Thus, in the absence of Barhl2, the majority of inhibitory Ptf1a:GFP cells remain in the appropriate layers. We next assessed the intrinsic effects of Barhl2 at single-cell level in a normal developing environment. For this we used the technical advantage of the zebrafish model to generate chimeras. Cells from embryos injected with Barhl2 MO or control MO were transplanted into wild-type host embryos (Figure 6A). We first transplanted cells from Tg(*ptf1a:GFP*) donors injected with *H2B-RFP* mRNA (to label all donor cells). Overall, the Ptf1a:GFP cells remained as inhibitory cells primarily in the inhibitory amacrine layer (22.96% in WT to 21.2% in morphants, p – value = 0.13, Figure 6B, C). The key question is whether the ACs

generated in *Barhl2* morphants show changed subtype identity. Using transplantations from WT donors combined with immunohistochemistry, we indeed found that amacrine subtypes that usually express *barhl2* were significantly lost in *Barhl2* morphant transplants (serotonin 43%, p – value = 0.04, calbindin 38%, p – value < 0.0001, GABA 68%, p – value < 0.0001, Figure 6D – G, J). In contrast, some subtypes that usually do not express *barhl2* were strikingly increased (neuropeptide Y 216%, p – value = 0.03), and others mildly increased (ChAT 125%, p – value = 0.14, Figure 6H – J). No evidence was found for a role of *Barhl2* in RGC differentiation or survival, as seen in other species (Mo et al., 2004; Poggi et al., 2005b; Ding et al., 2009). These observations demonstrate that the *Barhl2* paralog in zebrafish is uniquely dedicated to the AC fates, in which it biases precursors towards generating specific some AC subtypes and away from others. These results also highlight a correlation between the expression level of *barhl2* within distinct AC subtypes and its necessity during the development of each subtype.

Barhl2 ACs are nieces of RGCs

The results above demonstrate that *Barhl2* acts downstream of *Ptf1a* to bias specific subtype identities within a subset of post-mitotic AC precursors. How does this cell population and transcriptional cascade relate to the *Atoh7*-lineage *in vivo*? Approximately 2/3 of the AC population arises from *atoh7*-expressing progenitors (Jusuf et al., 2011). Since we found that *barhl2*-expressing neurons comprise 58.5% of *ptf1a*-expressing ACs, we wondered whether *barhl2*-expressing ACs come from *atoh7*-expressing progenitors. Previous expression analysis on *barhl2* and *atoh7* did not provide any evidence for mRNA co-localization, e.g. RGCs express *atoh7* and not *barhl2* and ACs that express *barhl2* do not express *atoh7* (Schuhmacher et al., 2011). This kind of approach, however, cannot rule out possible lineage-relationships between cells expressing these two factors. In contrast, some of the fluorescent proteins display long perdurance compared to the native mRNAs under whose promoters they are driven. Thus, in the *Tg(barhl2:GFP/atoh7:gap43-RFP)* double transgenic line in which the RFP is long-lived compared to *atoh7* mRNA, we can visualize the respective onset of *Barhl2:GFP* and *Atoh7:gap43-RFP* in individual cells using *in vivo* 3D time-lapse imaging. We started imaging at around 35 hpf, when retinal neurons first express *Barhl2:GFP*, and asked whether any *Barhl2:GFP*-positive neurons arise from *atoh7*-expressing progenitors within the first 6 hours of each time-lapse movie (Figure 7A). Strikingly, we found that almost all of the *Barhl2:GFP*-expressing cells in this time-window also were *Atoh7:gap43-RFP*-positives (94.3% \pm 2.8% SEM), suggesting that *Barhl2*-dependent amacrine subtypes indeed arise from the *Atoh7*-lineage. We compared the distribution of subtypes that express *barhl2* with our previously identified subtype biases within the *Atoh7*-lineage (Jusuf et al., 2011). We thus performed morphological characterisation of individual *Barhl2:GFP* expressing cells ($n = 28$, Figure 7B - D) and compared the frequency of subtypes in three categories: We found that *Barhl2:GFP* subtypes of ACs are underrepresented or overrepresented at similar frequency as are *Atoh7+*/*Ptf1a+* expressing cells when compared to all (*Ptf1a+*) subtype frequencies. Thus, our *Barhl2+* sample was statistically significantly different from WT (p – value = 0.032), but comparable with the *Atoh7+*/*Ptf1a+* population (p – value = 0.55), which our time-lapse analysis showed to be the origin of *barhl2*-expressing cells.

If *Barhl2*-dependent ACs derive from the *Atoh7*-lineage, then time-lapse imaging should allow us to understand the lineage relationship between *barhl2*-expressing amacrine subtypes and RGCs. In our previous 3D time-lapse study we traced individual dividing *Atoh7:GFP*-expressing progenitors long enough to show that these cells often divided asymmetrically to produce one daughter cell that became a RGC and another daughter which often migrated back toward the apical surface (Poggi et al., 2005b). The time-lapse limitations and lack of appropriate fluorescent reporters did not allow us to follow the fate of

the non-RGC daughter cell for more than a few hours, during which time they did not divide again (Poggi et al., 2005b). In the Tg(*barhl2:GFP/atoh7:gap43-RFP*) double transgenic line, however, it is possible to trace the cellular origin of Barhl2:GFP/Atoh7:gap43-RFP-positive cells. Seven retinas were imaged for a minimum of 20 hours starting from 28 hpf. We traced the lineage of 20 individual Barhl2:GFP/Atoh7:gap43-RFP positive cells. Strikingly, in 19 out of these 20 cases, the Barhl2:GFP positive cell arose from a cell division of an Atoh7:gap43-RFP-positive progenitor, which was identified as the sister of an RGC (Movie 2). In all these 19 cases, the Atoh7:gap43-RFP progenitor generated one Barhl2-positive and one Barhl2-negative daughter cell whose identity remains unknown (Movie 2). In one case out of the 20, we observed that both daughters became Barhl2:GFP-positive. In 14 cases, we were able to reconstruct the lineage of the Barhl2-positive AC from the very onset of Atoh7:gap43-RFP expression. In all of these cases, the lineage started with the asymmetrical division of an Atoh7-positive progenitor, one daughter of which turned up Atoh7:gap43-RFP expression and became a RGC, while the other daughter divided again (see example in Figure 7E). The result of this next division was one Barhl2:GFP-positive cell and one Barhl2:GFP-negative cell (Figure 7E and Movie 2). Taken together, these observations suggest that during RGC genesis *in vivo*, Barhl2-positive ACs arise as nieces of RGCs, mainly through asymmetric fate outcome of *atoh7*-expressing progenitors division (Figure 7F).

Barhl2 acts within the Atoh7-lineage to specify subtypes of ACs

Barhl2 is expressed in cells that derive from Atoh7-progenitors, and biases AC identities towards the same subtype identities that arise in the Atoh7-lineage. Therefore, it is important to determine whether Barhl2 by itself is capable in biasing amacrine subtypes *in vivo*. We investigated this by expressing *barhl2* within *atoh7*-expressing progenitors such that all rather than a subset of these cells would turn on Barhl2. DNA for the H2B-RFP reporter was injected into 1-cell stage embryos in which either *pUAS:H2B-RFP* (in control retinas) or from *pUAS:barhl2-T2A-H2B-RFP* (*barhl2* misexpressing retinas) was expressed under the control of *atoh7* promoter in the Tg(*atoh7:gal4/pUAS:gap43-GFP*) line. The use of Barhl2-T2A-H2B-RFP in frame fusion allows visualization of *barhl2*-expressing cells *in vivo*, and allows simultaneous assessment of their final fate (Kim et al., 2011). *In vivo* time-lapse analysis confirmed that the expression of *barhl2* (H2B-RFP positive cells) indeed occurred prematurely, in dividing Atoh7-positive progenitors (Movie 3). Cell fates of H2B-RFP labelled cells were quantified at 4 dpf (Figure 8). We found no significant changes in the proportion of H2B-RFP positive ACs in the INL ($21.02\% \pm 4.08$ SEM control to $26.9\% \pm 4.15$ SEM misexpression, p – value = 0.3255) or HCs ($20.26\% \pm 5.11$ SEM control to $19.26\% \pm 4.0$ SEM misexpression, p – value = 0.5157), nor bipolar cells in the outer half of the INL ($8.2\% \pm 5.02$ SEM control to $8.9\% \pm 3.97$ SEM misexpression, p – value = 0.1404). However, *barhl2* misexpression leads to a significant increase of H2B-RFP positive cells in the GCL ($17.68\% \pm 5.36$ SEM control to $34.82\% \pm 5.8$ SEM misexpression, p – value = 0.0017) and a significant loss of H2B-RFP positive cells amongst photoreceptor (the outer nuclear layer (ONL)) cells ($32.84\% \pm 5.76$ SEM control to $10.12\% \pm 5.4$ SEM misexpression, p – value = 0.0105). To analyse this effect with respect to AC subtypes, we used antibodies for specific AC populations. Within *barhl2*-misexpressing H2B-RFP positive cells we found a significant increase in the proportion of GABAergic cells in the INL ($15.93\% \pm 4.86$ SEM control to $22.2\% \pm 4.64$ SEM misexpression, p – value = 0.0044) and GCL ($7.5\% \pm 4$ SEM control to $28.32\% \pm 6.05$ SEM misexpression, p – value < 0.0001), which was particularly evident in the GCL (Figure 8D). Therefore, the increase in GABA+ cells appears to be at the expense of early made ganglion cells (GABA- in the GCL) and photoreceptors. Notably, neither in the control nor in the misexpression condition were H2B-RFP positive cells ever ChAT-positive (data not shown), a subtype that rarely expresses *barhl2*. These results suggest that when *barhl2* is prematurely expressed in the

atoh7-expressing progenitors population, it is itself able to induce within these cells some aspects of Barhl2-dependent AC subtype identities.

Atoh7 affects lineage outcome and number of Barhl2-positive cells

We know that *ptf1a* expression does not depend on Atoh7 (Jusuf et al., 2011), but as *barhl2* is turned on in Atoh7-derived inhibitory precursors, we wondered whether Atoh7 influences the way Barhl2-positive cells arise within the Atoh7-lineage. The Tg(*barhl2:GFP*) line was outcrossed to *atoh7^{-/-}* (*lakritz*) mutants (Kay et al., 2001). Barhl2:GFP expression analysed at 5 dpf is not only retained, but also virtually doubled in the absence of Atoh7 ($39.33\% \pm 0.92$ SEM WT to $73.74\% \pm 2.261$ SEM *lakritz*, p – value < 0.0001 ; data not shown) with significant increases in the INL ($48.33\% \pm 1.32$ SEM to $78.32\% \pm 2.35$ SEM, p – value < 0.0001 ; data not shown) and GCL ($20\% \pm 1.03$ SEM to $63.89\% \pm 3.81$ SEM, p – value < 0.0001 ; data not shown). How do Barhl2-positive ACs arise in this case? In twelve time-lapse movies of Barhl2:GFP/Atoh7:gap43-RFP cells in *lakritz* retinas, we consistently observe Atoh7:gap43-RFP-positive cell divisions generating two daughters, only one of which becomes Barhl2:GFP-positive. Interestingly, in eight of these movies, the original Atoh7:gap43-RFP cells first divide once as usual. However, as neither daughter is able to differentiate into a RGC, both daughters divide again, with one cell from each pair starting to express Barhl2:GFP (Movie 4). Thus, when Atoh7 is missing, Barhl2-positive ACs still arise within asymmetric cell divisions from the “Atoh7-lineage”, but in this case there are two rather than only one *barhl2*-expressing cell generated from each lineage. This different lineage-outcome reveals the mechanism of Barhl2-positive amacrine cell’s increased number in the *lakritz* mutant.

Atoh7 affects Barhl2-positive amacrine subtypes identity

Are *barhl2*-expressing subtypes increased equally in the absence of Atoh7? Since GABA labels the majority of Barhl2:GFP cells, we first assessed changes to this cell population. We found that the total proportion of Barhl2:GFP/GABAergic neurons remains unchanged in *lakritz* mutants (32% WT to 30% *lakritz*). Because GABAergic neurons consist of multiple subtypes, we next assessed, if specific subtypes are changed within this population. We found that serotonergic amacrine subtypes (usually *barhl2*-expressing) are significantly increased ($+20.6\% \pm 6$ SEM *lakritz* retina, p – value = 0.0002, Figure 9E – G), thus suggesting that specific *barhl2*-expressing subtypes are preferentially affected. The most striking change is the large increase in Barhl2:GFP non-GABAergic amacrine cells in the *lakritz* retina (Figure 9C, D). One possible explanation for this outcome could be that Barhl2-negative subtypes now become Barhl2-positive. To investigate on this possibility, we analysed the population of NY+ amacrine subtypes. We found no significant change in NY+ proportion or GFP expression ($+$ or $-$ $11\% \pm 5.5$ SEM *lakritz* retina, p – value = 0.2611). Our data thus show that specific subtypes of Barhl2:GFP+ cells, including 5-HT+ and non-GABAergic populations, are preferentially expanded. Although the increase in amacrine cells can be explained by our lineage analysis, the mechanism by which Atoh7 affects subtype identity remains unclear.

Lakritz mutants retain some cells in the ganglion cell layer of the retina, which have been attributed to displaced amacrine cells (Kay et al., 2001). Does this simply reflect the tendency of ACs to occupy the now RGC-free most basal positions in the retina (Kay et al., 2001), or rather it indicates a selective increase in specific subclasses of amacrine cells (Feng et al., 2010), e.g. the Barhl2:GFP cells deriving from the Atoh7-lineage? To gain more insights into this question we analysed GABA as marking a large proportion of the usually Barhl2+ population, as well as serotonin and neuropeptide Y as subtype specific markers that were usually Barhl2+ or Barhl2-respectively. We found GABAergic Barhl2:GFP increased in the GCL (12.8% WT to 21% *lakritz*, p -value = 0.0008) and

decreased in the INL (19.2% WT to 9% *lakritz*, p-value = 0.0014) (Figure 9A - D). Similarly, serotonergic *Barhl2*:GFP cells, which are normally only found in the INL (Figure 9E - G) are now increased due to their appearance in the GCL (+20.2% increase, p - value = 0.0002, Figure 9E - G). Finally, even though the proportion of NY+ remained unchanged in the *lakritz* retina, some of these cells were redistributed to the GCL (11% increase in the *lakritz* retina compared to the WT, p - value = 0.0009, **data not shown**). These observations together suggest that amacrine subtypes get redistributed to the ganglion cell layer in the *lakritz* mutants, regardless of whether they are re-specified or increased in number.

DISCUSSION

Although neural cell fate determination factors are being studied in various model organisms, the occurrence of particular patterns and interactions within individual cell lineages can only be assessed *in vivo* in the zebrafish model system. In this study we use a combination of functional and *in vivo* time-lapse analyses to uncover novel aspects of lineage-related transcriptional networks and cell-fate outcomes in the developing vertebrate retina. In particular, we here demonstrate that the zebrafish *Barhl2* paralog biases particular ACs *in vivo*, and is turned on in particular *Atoh7*-lineages.

In the zebrafish retina, 71% of inhibitory neurons (all horizontal and about 60% of amacrine cells) derive from *Atoh7* progenitors. These are distinct in subtype composition from the remaining third of amacrine subtypes that derive from non-*Atoh7* progenitors (Jusuf et al., 2011). Most amacrine subtypes can arise from either progenitor, but the probability of assuming a particular subtype fate correlates with whether the AC arose from an *Atoh7*-positive or -negative lineage. Experimentally induced co-expression of *ptf1a* in all *Atoh7*-positive progenitors leads to a selective increase in the ACs that normally derive from the *Atoh7*-lineage (Jusuf et al., 2011). Yet *Atoh7* is not essential for the specification of any of the amacrine subtypes, and *ptf1a* is expressed in all ACs, suggesting that other factors must be key to biasing subtype specification. Here we find that 58.5% of amacrine cells express *barhl2*, and that most *Barhl2* cells arise from *atoh7*-expressing progenitors, thus providing evidence that *Barhl2* may be this “other” factor. Consistent with this hypothesis, we found that *barhl2*-expressing subtypes were similarly underrepresented and overrepresented when compared to our previously characterised *Atoh7*+/*Ptf1a*+ lineage. Furthermore, knocking down or increasing *barhl2* expression within *Atoh7*-progenitors is sufficient to bias the differentiation of amacrine cells in predictable directions. Taken together these observations provide an explanation as to why the *ptf1a*-expressing progenitors in the *Atoh7*-lineage, but not those from the non-*Atoh7* lineage, bias differentiation toward specific amacrine subtypes and not others (Jusuf et al., 2011).

Barhl2 has been implicated in the development of ACs, and specification and survival of RGCs downstream of *atoh7* in the mouse and *Xenopus* retina (Mo et al., 2004; Poggi et al., 2004; Wang and Harris, 2005; Ding et al., 2009). Here we find that the zebrafish *barhl2* paralog is turned on exclusively in post-mitotic, inhibitory *Ptf1a*-positive ACs and horizontal cells. We also find that *barhl2* is preferentially expressed at different levels in specific amacrine subtypes (such as GABA, 5-HT and calbindin expressing), whereas it is almost completely absent in others (NY and ChAT). In accordance with data from mouse, the absence of *Barhl2* function leads to an altered amacrine subtype composition (Ding et al., 2009). Specifically, subtypes that are normally *Barhl2*-positive (GABA, serotonin and calbindin) decrease at the expense of the ones that normally do not express *barhl2* (ChAT and NY). Decreases in GABAergic population with increases in cholinergic AC are consistent with the previous mouse data (Ding et al., 2009) and these may indeed represent homologous subtypes. However, other subtypes may present preferential alternate fates, as the proportional increase in NY cells far outweighed the modest and statistically not

significant increase in ChAT⁺ cells. Although some of the markers may label overlapping amacrine subtype populations, our results show specific losses in subtypes that express *barhl2* most strongly. We therefore show for the first time a strong direct correlation between expression levels of *barhl2* and cell-autonomous necessity for Barhl2 in each of these specific subtypes, rather than just a general role in relative subtype composition.

Unlike in other vertebrates (Poggi et al., 2004; Ding et al., 2009), the zebrafish *barhl2* is not likely implicated in RGC differentiation or survival. Several observations support this conclusion: (1) *barhl2* expression is restricted to *ptf1a*-expressing inhibitory cell precursors and is turned on only after the division of sister cell of an RGC; (2) Overexpression of *barhl2* is sufficient for amacrine subtypes, but not RGCs, even if overexpressed prematurely under the control of the *atoh7* promoter; (3) knocking down *barhl2* does not show significant changes in RGC genesis; (4) *barhl2* expression depends on the expression of *ptf1a* and not on the expression of *atoh7*. All of these observations are in agreement with Barhl2 acting exclusively in inhibitory cell precursors. The function of *barhl* genes in RGCs genesis or maintenance might instead be retained by the duplicated *barhl1.2* paralog; which is expressed in RGCs downstream of *Atoh7* (Schuhmacher et al., 2011).

What is then the relationship between *barhl2* and *atoh7*? Previous expression analysis suggested that a negative feedback might occur between the two genes, possibly as a result of the conservation of an ancestral feature existing in the invertebrate *Drosophila melanogaster* between the homologues *atonal* and *barH* genes (Lim and Choi, 2003; Schuhmacher et al., 2011). Our *in vivo* timelapse study highlighted particular lineage-relationships between *Atoh7*-dependent RGCs and *barhl2*-expressing amacrine subtypes. Our results clearly demonstrate that the expression of the two genes is mutually exclusive at cellular level; they also show that Barhl2-positive ACs arise as nieces of RGCs, which depend on *Atoh7*. This observation, combined with results from loss- and gain-of-function assigns the timing of Barhl2 action in post-mitotic precursors in which *atoh7* has been already down-regulated (Skowronska-Krawczyk et al., 2005; Le et al., 2006; Brzezinski et al., 2012) and the switch from RGC to inhibitory cell fates has already occurred *via* the expression of *ptf1a*. Thus, Barhl2 *in vivo* is not well positioned to act as a feedback repressor of *atoh7*. The intermediate *ptf1a* expression however, might in fact repress *atoh7* within this lineage consistent with data in chick (Lelievre et al., 2011) (Figure 7F).

Conversely, we found that *barhl2* expression does not need functional *Atoh7*. *Atoh7* might therefore exert an indirect repressive function on *barhl2*. A recent study in the mouse shows that in *Math5/atoh7*-null retina there is a precocious expression of AC fate determinants, thus suggesting that *Atoh7* prevents ACs from being generated prematurely (Feng et al., 2010). Yet our *in vivo* time-lapse analysis of *barhl2:GFP/atoh7:gap43-RFP* progenitors through two rounds of cell division shows that the onset of Barhl2:GFP within individual cell lineages is unvaried in the absence of *Atoh7*. Consistently, in both wild type and *lakritz* retina, Barhl2-dependent ACs tend to arise from divisions of “*Atoh7:gap43-RFP*” cells in which one daughter turns on Barhl2:GFP and the other does not. However, we also find that *Atoh7* affects the lineage-outcome of retinal progenitor cells. Therefore, the increase in the number of Barhl2-positive cells is rather due to the fact that without *Atoh7*, both daughters of one *Atoh7:gap43-RFP* progenitor divide again to generate an additional Barhl2-positive daughter. This is coherent with a co-temporaneous study by He et al. (2012 in press), in which randomly selected lineages in the zebrafish retina were traced in time-lapse. In these lineages, approximately 80% of all RGCs arise from asymmetric divisions in which one daughter differentiates as an RGC and the other daughter divides again. Results from He et al. also show that in the absence of *Atoh7*, there is a marked decrease in such divisions.

Although the onset of *barhl2* is independent of *Atoh7* action, our results clearly show that *Atoh7* does affect *barhl2*-expressing amacrine subtypes identity. While GABAergic/*Barhl2*-positive amacrine cells remain unchanged in the *lakritz* retina, there is a striking increase in the non-GABAergic/*Barhl2*-positive cells and in the normally rare serotonin/*Barhl2*-positive ACs. Thus, *Atoh7* might be permissive for some *Barhl2*-dependent amacrine subtypes (such as GABA) while it might repress others (e.g. serotonin). In the *atoh7*-null retinas of the mouse, a change in amacrine subtype composition has also been noted, with certain amacrine subtypes increasing at the expense of others (Feng et al., 2010). Also in agreement with these studies, our analysis of the *lakritz* retina shows an overall increase in displaced ACs (Kay et al., 2001; Feng et al., 2010). This general increase was happening regardless of whether ACs were particular *Barhl2*-positive or *Barhl2*-negative (such as for NY) subtypes.

It remains a major challenge to identify cell fate determination networks and how they operate within the context of lineage in the developing vertebrate CNS. Recent studies suggest that there are stochastic mechanisms at work in determining fate choice within clones (Gomes et al., 2011; He et al., 2012 in press), and perhaps that is why factors like *Barhl2* bias rather than strictly determine fate choice, but our results showing the rather strict expression of *Barhl2* within the *Atoh7*-lineage argue that the expression of at least some cellular determinants in the retina are strongly influenced lineage. We have shown that the zebrafish retina is an excellent model system to address this challenge as this system allows us to follow *in vivo* modes of cell divisions within individual lineages, concurrently with the expression of specific cell fate determinants. Here, we highlighted the lineage of individual *atoh7*-expressing progenitors through two rounds of cell divisions, and our functional results suggest that interactions involving the division mode, and the transcriptional network of *Atoh7*, *Ptf1a* and *Barhl2* promote the orderly generation of RGCs and amacrine subtypes within this lineage.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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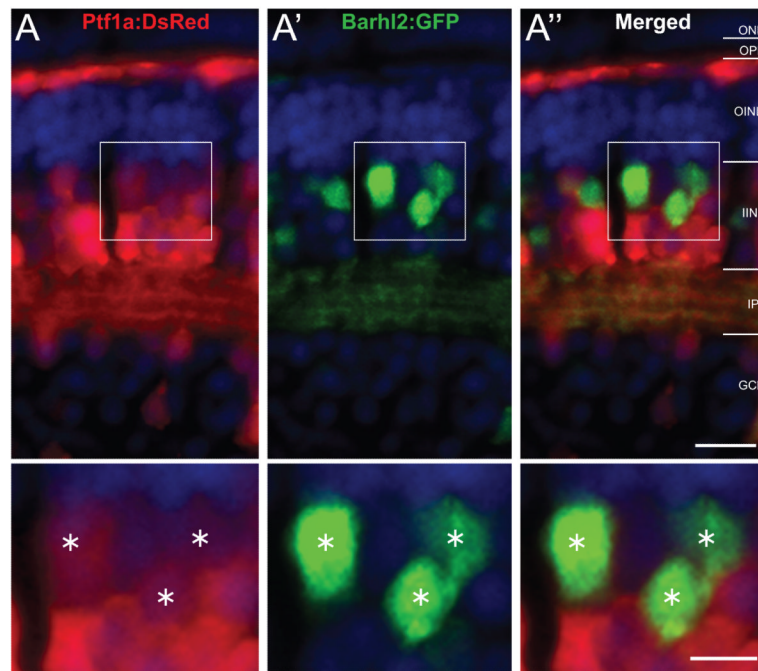
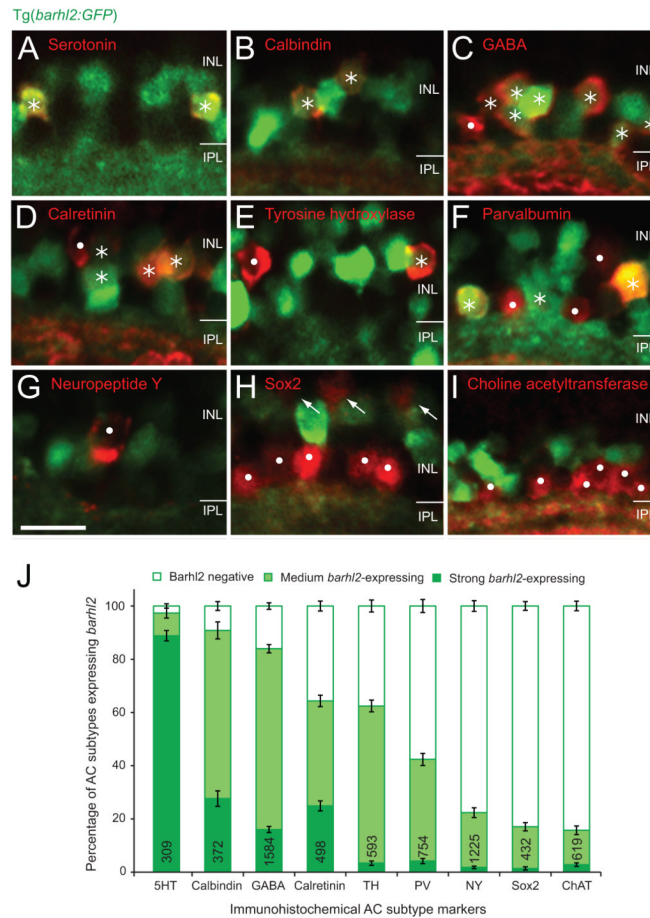


Figure 1.

Barhl2:GFP labels a subpopulation of Ptf1a:DsRed cells. Micrographs of retinal sections from *Tg(ptf1a:dsRed/barhl2:GFP)*. Cell nuclei (in blue) are stained with DAPI. All of the Barhl2-positive cells are positive for varying degrees of Ptf1a. In contrast, only some of the DsRed cells also express GFP, showing that a subpopulation of *ptf1a*-expressing cells expresses *barhl2*. Most of these double-labelled cells are found in the amacrine population in the INL. Results quantified from $n = 58$ eyes, 5358 cells in central retina of 96 hpf embryos. ONL: outer nuclear layer; OPL: outer plexiform layer; OINL: outer half of the inner nuclear layer; IINL: inner half of the inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell layer. Asterisks mark cells expressing both *GFP* and *dsRed*. Scale bar A'' (for A – A'') = 20 μm , scale bar A'' inset (for A – A'' insets) = 10 μm .

**Figure 2.**

Amacrine markers reveal distinct subtypes expressing *barhl2*. Micrographs of 120 hpf *Tg(barhl2:GFP)* embryos immunohistochemically labelled with markers (red). (A – C) Serotonin (5HT, A), calbindin (B) and GABA (C) immunoreactive AC types primarily co-label with *Barhl2:GFP*. (D – F) Calretinin (D), tyrosine hydroxylase (TH, E) and parvalbumin (PV, F) labelled populations show varying degree and intensities of *Barhl2:GFP* co-labelling. (G – I) Neuropeptide Y (NY, G), Sox2 (H) and choline acetyltransferase (ChAT, I) label ACs that do not co-label with *Barhl2:GFP*. Sox2 additionally labels Müller glia cells (arrows, H). Asterisks indicate ACs (red) that co-localise with GFP; dots mark cells that do not express GFP. (J) Quantification of the percentage of different markers co-labelled with *Tg(barhl2:GFP)*, $n = 21 - 220$ eyes, 309 – 1584 cells. ACs labelled by 5-HT, calbindin and GABA primarily arise from *barhl2*-expressing cells. ACs labelled by NY, Sox2 and ChAT primarily come from *Barhl2*-negative or weakly *barhl2*-expressing cells. ACs labelled with calretinin, TH or PV include both, cells that do and do not arise from *barhl2*-expressing cells. AC: amacrine cell. Numbers in each bar indicate the number of labelled cells analyzed. Error bars indicate standard error of the mean. INL: inner nuclear layer; IPL: inner plexiform layer. Scale bar G (for A – I) = 20 μm .

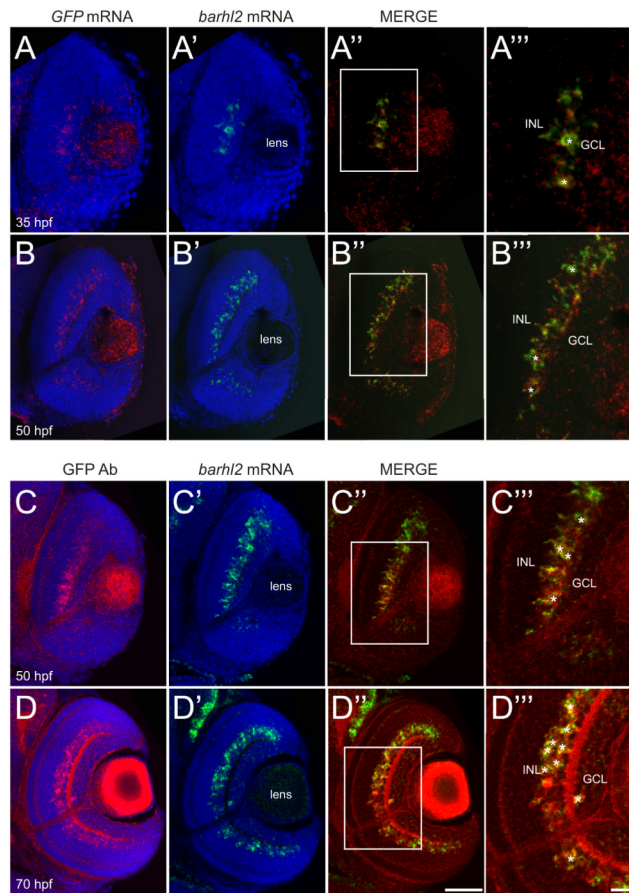


Figure 3.

The Tg(*barhl2:GFP*) transgenic line faithfully reflects the endogenous *barhl2* expression in time and space. (A – B) Whole-mount double fluorescent *in situ* hybridization against *barhl2* mRNA and *GFP* mRNA on zebrafish retinas counterstained with DAPI (blue) at 35 hpf (A – A'') and 50 hpf (B – B''). (C – D) Whole-mount fluorescent *in situ* hybridization against *barhl2* mRNA followed by immunohistochemical labelling against GFP at 50 hpf (C – C'') and 70 hpf (D – D''). Co-localization of *barhl2* mRNA with *barhl2* mRNA or Barhl2:GFP expression (asterisks) occurs in the inner nuclear layer (INL) at all ages and in few cells in the ganglion cell layer (GCL) at 70 hpf. Scale bar D'' (for A – D'') = 50 μ m, scale bar D''' (for A''' – D''') = 20 μ m.

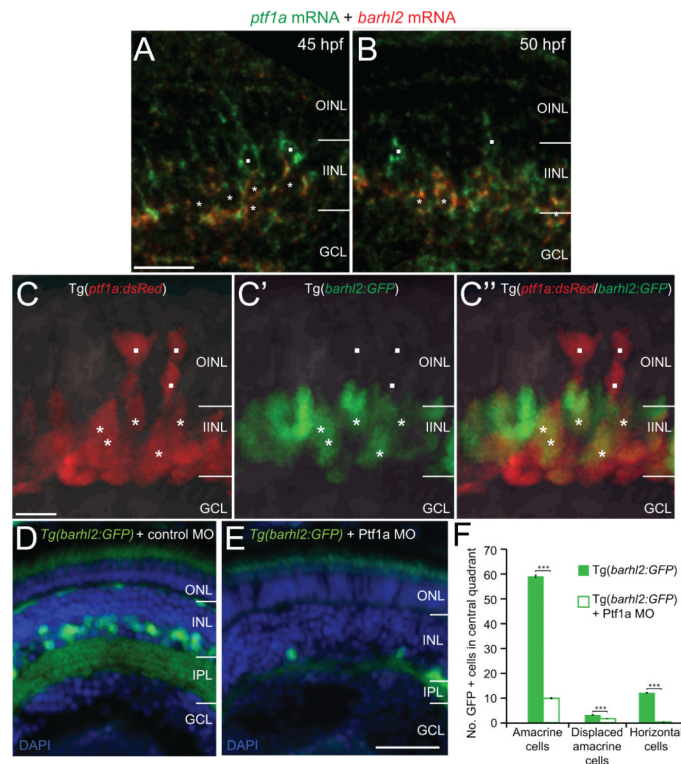


Figure 4.

Ptf1a / *barhl2* are sequentially expressed within individual cells, with *Ptf1a* being necessary for *Barhl2:GFP* expression. (A, B) Double fluorescent *in situ* hybridization of *barhl2* and *ptf1a* mRNAs. *Ptf1a* (green) is expressed in cells apically (squares) and in cells that have migrated basally where they also express *barhl2* (red, asterisks) at 45 hpf (A) and 50 hpf (B). (C – C'') Micrographs from *in vivo* time-lapse of double transgenic Tg(*ptf1a:dsRed*/*barhl2:GFP*) embryos at 35 hpf show a similar pattern with apical cells expressing *Ptf1a:DsRed* alone (squares) and more basal cells co-expressing *Barhl2:GFP* (asterisks). (D, E) Micrographs of 120 hpf Tg(*barhl2:GFP*) injected with standard morpholino (D) or *Ptf1a* morpholino (E). *Barhl2:GFP* expression is drastically reduced in the *Ptf1a* morphants. (F) Quantification of GFP labelled cells shows a significant loss of cells in *Ptf1a* MO in all cell types that usually express *Barhl2:GFP*, i.e. ACs (inner half of INL), displaced ACs (outermost layer of GCL) and horizontal cells (outermost layer of INL). WT n = 51 eyes, *Ptf1a* morphant n = 41 eyes. ONL: outer nuclear layer; INL: inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell layer; MO: morpholino; ***: p – value < 0.001; error bars indicate standard error of the mean. Scale bar A = 20 μ m (for A, B), scale bar C (for C – C'') = 10 μ m, scale bar E (for D, E) = 50 μ m.

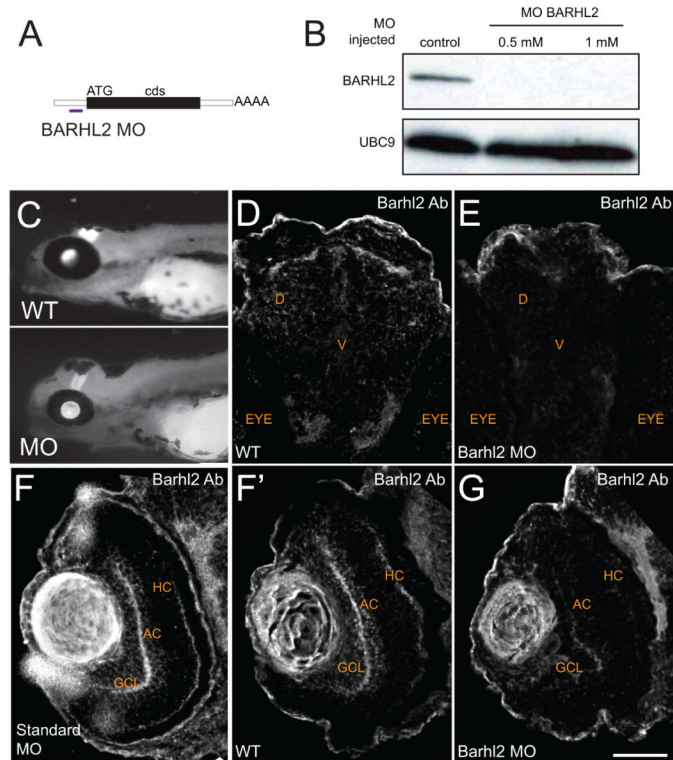


Figure 5. Zebrafish *Barhl2* morpholino causes efficient *barhl2* knockdown. **(A)** A translational blocking morpholino (MO) was designed against 6 base pairs upstream the translational start site of zebrafish *barhl2* mRNA. The morpholino efficiency was tested using anti-*Barhl2* antibody. **(B)** Western Blotting of wild type (control) and morphant (0.5 and 1 mM *Barhl2* MO) 50 hpf old embryos heads. **(C)** Micrographs of control and *Barhl2* morphant embryos. Morphants look relatively normal, showing only mild general defects in their heads. **(D – G)** Immunohistochemical labelling of brain (D, E) and retinal (F, F', G) sections at 50 hpf. Control embryos (D, F) and standard MO injected embryos (F') show *Barhl2* protein in the diencephalon region D around the ventricular zone V of the brain (D) and in the retina (F, F'), in amacrine and horizontal cells (AC, HC) and in cells located in the ganglion cell layer (GCL, F, F'). *Barhl2* morphant embryo brain and retina do not stain for *Barhl2* protein. Scale bar G (for D – G) = 50 μ m.

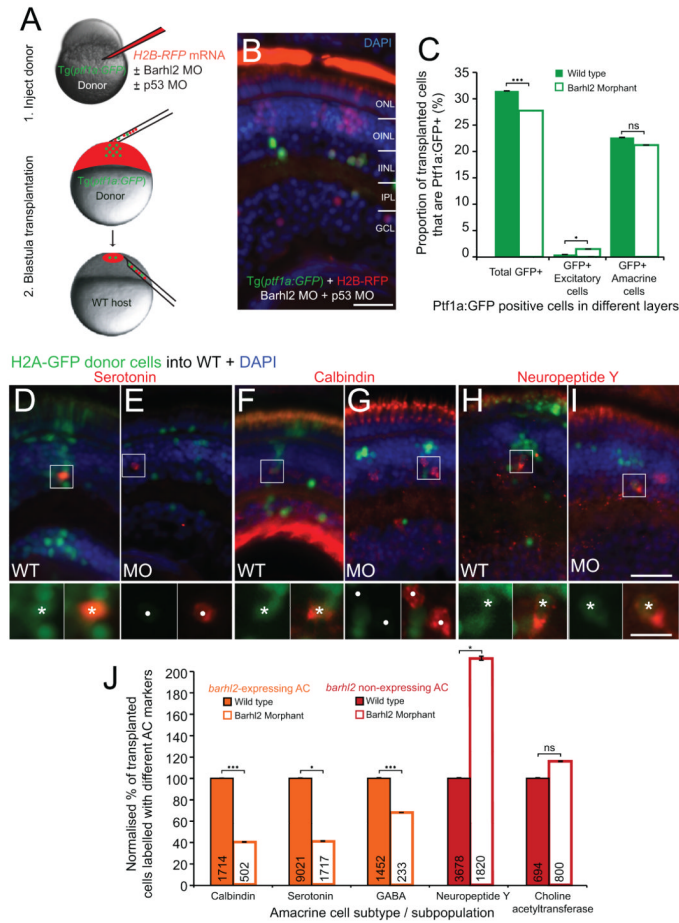


Figure 6. Barhl2 knockdown causes a subtype fate switch towards alternate inhibitory subtypes. (A) The cell-autonomous effect of Barhl2 loss was assessed with transplantations: Donor *Tg(ptf1a:GFP)* embryos were injected with *H2B-RFP mRNA* and p53 MO (to aide cell survival (Robu et al., 2007)), with (knockdown) or without (control) Barhl2 MO. Donor cells were subsequently transplanted into unlabelled WT. (B) Some transplanted H2B-RFP cells express Ptf1a:GFP in both conditions. (C) Quantification reveals a small reduction in the proportion of Ptf1a:GFP labelled donor cells (32% WT to 27.8% morphant, p – value < 0.001) and a few mislocalised Ptf1a:GFP cells in excitatory layers (0.74% WT to 1.49% morphants, p – value = 0.012). Overall, the vast majority of Ptf1a:GFP remain as inhibitory ACs (22.96% WT to 21.2% morphants, p – value = 0.13). (D – I) Immunohistochemically labelled amacrine subtypes (red) in chimeric retinas arise from transplanted H2A-GFP labelled donor (e.g. asterisks for serotonin - D, calbindin - F, neuropeptide Y-H, I) and from unlabelled host cells (circles). (J) Quantification of the proportion of labelled transplanted cells shows varying degrees of loss in subtypes that usually express *barhl2* (orange) and increases in AC subtypes that usually do not (red). ONL: outer nuclear layer; OINL: outer half of the inner nuclear layer; IINL: inner half of the inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell layer; MO: morphant / morpholino; ns: not significant; *: p – value < 0.05; ***: p – value < 0.001; error bars indicate standard error of the mean. Scale bar B = 20 μ m, scale bar I (for D – I) = 25 μ m, scale bar I inset (for D – I insets) = 10 μ m.

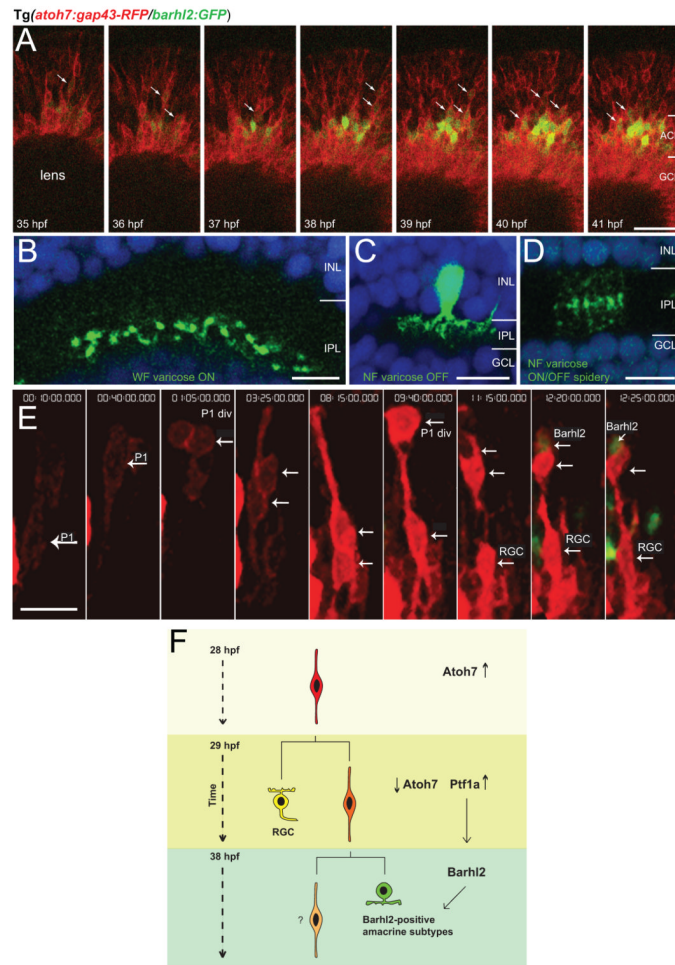


Figure 7.

Barhl2 is expressed in nieces of *atoh7*-expressing progenitors. (A) Micrographs from time-lapse movie ($n = 7$ movies) of double *Tg(atoh7:gap43-RFP/barhl2:GFP)* from 35 hpf. *Barhl2:GFP* cells almost always express *Atoh7:gap43-RFP*. *Barhl2:GFP* expression first occurs in cells that are migrating towards the amacrine layer. Boundaries are indicated for the forming amacrine (ACL) and ganglion cell layer (GCL), which contains the brightly *Atoh7:gap43-RFP* positive retinal ganglion cells (RGC). Arrows indicate co-labelling. *Barhl2:GFP* were analysed from the first frame in which they appeared towards the centre of the imaged stack (arrows). (B – D) Examples of individual amacrine cells arising from *barhl2* expression. Only one image of a confocal stack is shown, somas for some B and D were located at different depth. (E) *Barhl2*-positive cells derive from asymmetric divisions of RGC sisters in *atoh7*-expressing progenitors. 3D reconstruction of confocal stack from time-lapse series (starting at 28 hpf $t = 0h:00min$) shows the lineage of an *Atoh7:gap43-RFP* progenitor (P1). After P1 division (white arrow, P1 div, $t = 1h:05min$), both daughter cells migrate basally (white arrows, $t = 3h25min$). At $t = 8h15min$, one daughter cell migrates apically, dividing at $t = 9h40min$ (white arrow, P2 div), whereas the other daughter cell remains close to the basal surface, differentiating as an RGC. At $t = 12h20min$ one of the daughter cells from the P2 division up-regulates *Barhl2:GFP*. (F) Scheme summarising the observed lineage tree and sequence of genes expression. Time is indicated on the left. Arrows indicate *atoh7* being downregulated in the differentiating RGC and daughter cell while *ptf1a* is upregulated in the AC precursor (Jusuf and Harris, 2009; Jusuf et al., 2011;

Brzezinski et al., 2012). NF: narrow-field; WF: wide-field; P1: progenitor 1; P1 div: cell division of progenitor 1; P2 div: cell division of progenitor 2. Scale bar A = 30 μm , scale bars B – D = 10 μm , scale bar E = 7 μm .

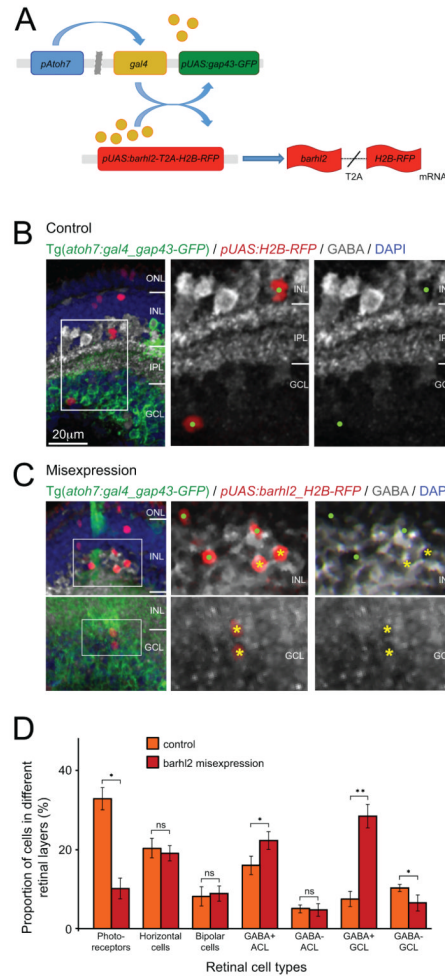


Figure 8.

Misexpression of *barhl2* in *atoh7*-expressing progenitors drives the fate of amacrine subtypes that usually express *barhl2*. (A) Schematic showing misexpression design: The *atoh7* promoter drives expression of Gal4 transcription factor. Gal4 activates the upstream activation sequence (pUAS) promoter to drive expression of gap43-GFP reporter by itself (control) or with Barhl2 and H2B-RFP reporter generated through in frame fusion with T2A peptide. After translation of the *barhl2-T2A-H2B-RFP* mRNA the T2A sequence will be cleaved and generate separate Barhl2 and H2B-RFP proteins. (B, C) Micrographs of 120 hpf *Tg(atoh7:gal4/pUAS:gap43-GFP)* retinas with pUAS driving *H2B-RFP* (control) or *barhl2* and *H2B-RFP* (misexpression), and subsequently labelled for GABA (white). Boxes indicate higher power insets. Cell type and GABA colabeling of *H2B-RFP*-expressing cells was analysed. Examples of GABA-positive cells (yellow asterisks) and GABA-negative cells (green dots). (D) Quantification of cell fates in control (orange) or *barhl2* misexpression (red). There is an increase in GABAergic cells in the INL (15.93% ± 4.86 SEM control to 22.2% ± 4.64 SEM misexpression, p – value = 0.0044) and particularly GCL (7.5% ± 4 SEM control to 28.32% ± 6.05 SEM misexpression, p – value < 0.0001 in GCL) with a concurrent loss in mainly photoreceptors (32.84% ± 5.76 SEM control to 10.12% ± 5.4 SEM misexpression, p – value = 0.01058) and presumed ganglion cells. ONL: outer nuclear layer; INL: inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell

layer; ns: not significant. Error bars indicate standard error of the mean. * p – value < 0.05 , ** p – value < 0.001 . Scale bar in B (for B, C) = 20 μm .

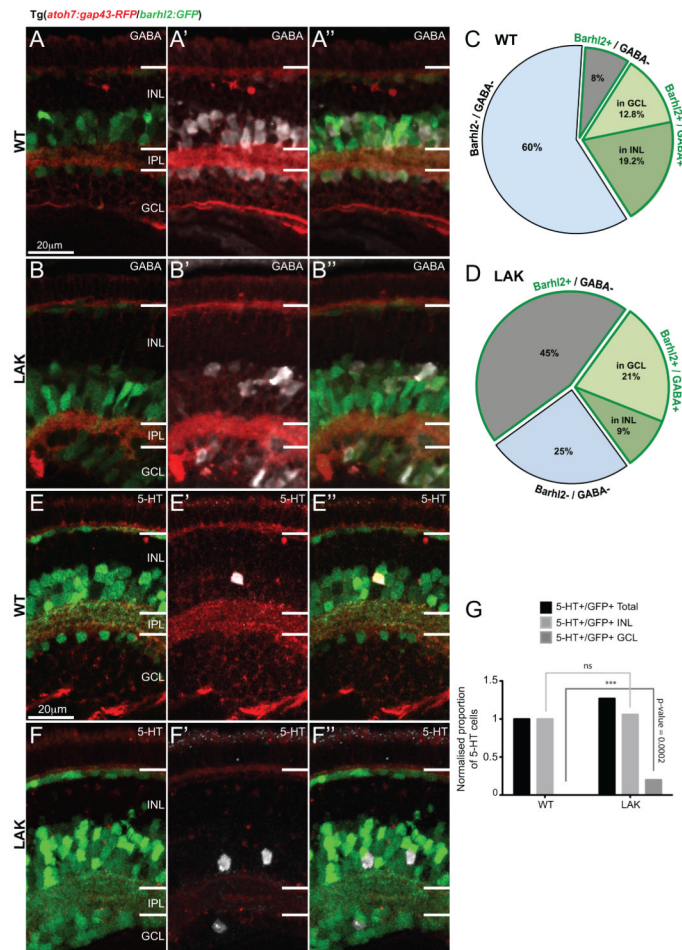


Figure 9.

Absence of *Atoh7* leads to an increase of *barhl2*-expressing ACs in the ganglion cell layer. (A, B) Micrographs of 120 hpf *Tg(atoh7:gap43-RFP/barhl2:GFP)* retinas labelled for GABA in wild type (WT) (A) and *lakritz* (LAK) (B). (C, D) Quantification of amacrine subpopulations expressed as proportion of total INL and GCL cells. The data indicate that the increase in the number of GFP+ cells in the LAK retina is primarily due to non-GABAergic ACs, as the proportion of GABAergic ACs remains unchanged in total in both GCL and INL in LAK vs. WT (32% WT to 30% *lakritz*). A redistribution of GABAergic ACs is however observed with an increase of Barhl2⁺/GABA⁺ cells in the GCL and a relative decrease of those cells in the INL of the LAK retina in comparison to WT (12.8% WT to 21% LAK, p-value = 0.0008 in GCL and 19.2% WT to 9% LAK, p-value = 0.0014 in INL). Total number of cells is the number of nuclei in the INL and GCL. (E, F) Micrographs of 120 hpf *Tg(atoh7:gap43-RFP/barhl2:GFP)* retinas labelled for serotonin (5-HT). (G) Quantification of the total proportion of serotonin-positive/Barhl2:GFP-positive (5-HT⁺/GFP⁺) neurons shows an increase in 5-HT cells in *lakritz* specifically in the GCL where they are normally absent (n = 15 – 19 eyes). IPL: inner plexiform layer; INL: inner nuclear layer; GCL: ganglion cell layer; *** p-value < 0.0002. Error bars indicate standard error of the mean. Scale bars = 20 μm.