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## Loss of Nance-Horan Syndrome b (*nhsb*) prevents expansion growth of retinal progenitor cells by selective up-regulation of $\Delta 113p53$ .

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Short Title: NHS represses  $\Delta$ 113p53 for retinogenesis

#### Abstract

The regulation of cell cycle progression and differentiation in retinal progenitor cells is a fundamental feature controlling organ size of the eye in vertebrates. Nance-Horan Syndrome (NHS) is a rare X-linked disorder caused by mutations in the *NHS* gene. Dysmorphic features of NHS include severe congenital cataracts, micropthalmia, facial dysmorphisms, and visual impairment. In this study we report an evolutionarily conserved role for NHS in vertebrate retinogenesis. Loss of function of *nhs* leads to small eye size in both zebrafish and *Xenopus tropicalis*, marked by reduced proliferation but not cell death. Transcriptome analysis of *nhs* morphant zebrafish eyes revealed a marked upregulation in  $\Delta 113p53$ , an isoform of *p53*, concomitant with a selective upregulation of p53 responsive genes that inhibit cell cycle progression but not apoptosis. Our data supports a model where Nhs is a negative regulator of  $\Delta 113p53$  expression and exerts its function through regulation of the p53 pathway to promote expansive growth of retinal progenitor cells prior to differentiation.

#### Keywords:

Nance-Horan Syndrome, nhsb,  $\Delta$ 113p53, p53, retinogenesis, zebrafish

#### Introduction:

A fundamental question in biology is how organ size is controlled during development. Organogenesis is regulated by a balance between progenitor cell cycle progression, self-renewal, differentiation, and programmed cell death. The neural retina serves as a model system that illustrates this general concept. The retina develops as an outpouching of the neuroepithelium and grows through symmetric proliferative divisions of retinal progenitor cells (RPCs) followed by a switch to asymmetric neurogenic divisions (Fadool and Dowling 2008; Willardsen and Link 2011; He et al. 2012; Boije et al. 2014). In the zebrafish retina, early RPCs undergoing symmetric cell divisions progress through the cell cycle slowly but then switch to a rapidly proliferating phase in which the cell cycle duration markedly decreases (Hu and Easter 1999; Li et al. 2000; He et al. 2012). This serves to rapidly increase the progenitor pool in advance of asymmetric neurogenic divisions in which some daughter cells exit the cell cycle to differentiate into neurons (He et al. 2012). Thus, coordinated cell proliferation and differentiation result in the correct number of cells that collectively contribute to the volume and size of the retina (Joseph and Hermanson 2010; Hindley and Philpott 2012). However, the molecular mechanisms that regulate cell cycle progression in retinal progenitor cells are not completely understood (He et al. 2012; Boije et al. 2014).

Nance-Horan Syndrome (NHS) is an X-linked developmental disorder characterized by severe congenital cataracts, dental anomalies, dysmorphic facial features and mental retardation (Brooks et al. 2004; Ramprasad et al. 2005; Coccia et al. 2009). Additional features of the disease include micropthalmia and microcornea (Florijn et al. 2006; Huang et al. 2007). NHS is caused by mutations in the *NHS* gene

found at the distal end of the X chromosome. The most frequent mutations associated with NHS are nonsense mutations predicted to result in a truncated protein (Burdon et al. 2003; Ramprasad et al. 2005; Florijn et al. 2006; Huang et al. 2007; Sharma et al. 2008; Coccia et al. 2009; Chograni et al. 2011; Tug et al. 2013; Hong et al. 2014; Shoshany et al. 2017; Tian et al. 2017), but insertions, small deletions, and a missense mutations have also been reported (Accogli et al., 2017; Brooks et al., 2004; Burdon et al., 2003; Chograni et al., 2011; Florijn et al., 2006; Li et al., 2015). Affected males have severe bilateral congenital dense nuclear cataracts that lead to profound vision loss and usually require surgery at an early age (Burdon et al. 2003; Huang et al. 2007; Sharma et al. 2008; Coccia et al. 2009). Despite cataract surgery during infancy, many NHS patients continue to exhibit visual impairment that may be caused by an underlying retinopathy (Mathys et al. 2007; Ding et al. 2009). It has been demonstrated that NHS protein is expressed in midbrain, lens, tooth and retina (Burdon et al. 2003; Sharma et al. 2006).

The p53 tumor suppressor protein has an established role as a cellular stress sensor, responding to a variety of insults such as DNA damage, nutrient starvation, nucleolar and ribosomal stress, and hypoxia (Vousden and Lane 2007; McCubrey et al. 2017). Evidence suggests that p53 functions to arrest cell cycle progression allowing time for DNA repair machineries to restore genome integrity before replication. If deleterious conditions are too severe, p53 activates cell death pathways. p53 performs many of its functions though downstream transcriptional activation of target genes, such as *cdkn1a/p21<sup>CIP1/WAF1</sup>* (p21) to halt cell cycle at the G1 phase and *bax* to execute programmed cell death pathways (Vogelstein et al. 2000; Harris and Levine 2005).

In addition to the full-length p53 protein, the p53 gene locus encodes for several isoforms generated by alternative promoter usage or alternative spicing (Bourdon et al. 2005; Joruiz and Bourdon 2016). A prominent variant  $\Delta 133p53$  in humans is transcribed from an internal promoter in intron 4 and is evolutionarily conserved in zebrafish (termed  $\Delta 113p53$ ), suggesting an important role in p53 functions (Bourdon et al. 2005; Chen et al. 2005).  $\Delta 113p53$  encodes an N-terminally truncated protein with a deletion of the transactivation domains and a partial deletion of the DNA-binding domain (Bourdon et al. 2005; Chen et al. 2005; Chen and Peng 2009).  $\Delta 113p53$  is itself a p53 target gene that is strongly upregulated following DNA damage, due to binding of full-length p53 to p53 response elements within intron4 of the p53 locus (Bourdon et al. 2005; Chen et al. 2005; Marcel et al. 2010; Aoubala et al. 2011). Previous work showed that Δ113p53 functions to protect cells from apoptosis by differentially modulating p53 response genes, promoting the expression of cell cycle arrest genes while antagonizing p53regulated apoptotic activity (Chen et al. 2009; Aoubala et al. 2011; Ou et al. 2014). Thus, the  $\Delta 113p53/\Delta 133p53$  variants function evolutionarily to promote cell survival when genome integrity is compromised to allow for DNA repair to take place (Gong et al. 2015).

Although p53 has a well-established role in maintaining genome stability to prevent tumor progression, a role for the p53 pathway in regulating development and organogenesis is less well understood. Loss-of-function mutations in p53 at first appeared to result mostly in viable offspring with no obvious developmental defects (Donehower et al. 1992). However, a number of p53 mutant mice die in utero due to exencephaly, a defect in neural tube closure due to overgrowth of neural progenitors in the midbrain, due at least in part to decreased apoptosis of neural progenitors (Sah et al., 1995). On the other hand, unrestrained activity of p53 can inappropriately induce p53 target genes and trigger cell cycle arrest and/or cell death during development (Jacobs et al. 2004; Dugani et al. 2009). For instance, loss of mdm2 or mdm4, negative regulators of the p53 protein, cause early embryonic lethality in a p53-dependent manner (Marine et al. 2006). More modest levels of p53 activation are associated with defects similar to those seen in patients with CHARGE syndrome (Van Nostrand et al. 2014). Recent work in zebrafish identified digestive organ expansion factor (def) as a pan-endodermal-enriched factor that is essential for expansive growth of digestive organs (Chen et al. 2005). Analysis of *def* null mutants revealed a selective regulation of the p53 isoform  $\Delta$ 113p53, that contributed to cell cycle arrest leading to reduced size and defective digestive organ development (Chen et al. 2005, 2009).

Here, we report a role for NHS in the development of the vertebrate retina. Knockdown of *nhs* in zebrafish and *frog* resulted in a reduced eye size. We show the small retinal size caused by *nhs* deficiency is due a decrease in RPC proliferation. The small retina, caused by nhs deficiency, is due a proliferation restriction mediated by the elevated expression of  $\Delta 113p53$ . We hypothesize that *nhs* functions in RPCs to limit p53 pathway induction as neural progenitors undergo rapid cell division.

#### RESULTS

#### The NHS family of genes

The Nance-Horan Syndrome family of genes is conserved among vertebrates including humans, mice, frogs, and zebrafish. Members of the NHS gene family include the prototypical NHS, as well as NHS-like 1 (NHSL1) and NHS-like 2 (NHSL2). The NHS genes are orthologs of the *Drosophila* gene GUK-holder (gukh)(Katoh and Katoh 2004)(Fig. 1A). Although most vertebrates express three *NHS* paralogs, (*NHS*, *NHSL1* and *NHSL2*), zebrafish have two copies of each paralog (*nhsa*, *nhsb*, *nhsl1a*, *nhsl1b*, *nhsl2a*, and *nhsl2b*) due to a genome duplication event in the teleost lineage (Postlethwait et al. 1998, 2000)(Fig. 1A).

In this study, we focused only on NHS orthologs. Previously, it was noted that the human NHS gene encompassed an unusually long distance between exon1 and exon2 (Brooks et al. 2004, 2010b). Comparison of other vertebrate genomes showed that the genomic architecture of the NHS gene remained conserved among human, mouse, frog and zebrafish with comparable number of exons. The large distance between exon1 and exon2 of NHS was also detected in other vertebrate genomes (Fig. 1B). The human NHS gene generates multiple protein isoforms by virtue of alternative promoter usage and alternative splicing (Huang et al. 2006; Brooks et al. 2010a). Isoforms arising from the alternative translational start site in exon1a appear to be evolutionarily conserved across vertebrates, as evidenced by sequencing of expressed sequence tags (EST) libraries (Fig. 1B). However, only NHS isoforms encoded from the first translational start

site in exon1 are predicted to be essential for development, since nonsense mutations in exon1 in humans leads to NHS (Burdon et al. 2003; Brooks et al. 2004).

We examined the temporal expression pattern of *nhs* paralogs during zebrafish development and find that both *nhsa* and *nhsb* are present in embryos before 2 hours post fertilization (hpf), suggesting that both transcripts are loaded maternally (Fig. 1C). We find that both *nhs* paralogs are expressed through all stages of development (Fig. 1C).



## Zebrafish *nhsb* is expressed in the developing eye and is required for normal retinogenesis

To determine the role of NHS in zebrafish development, we first examined the *nhsa*<sup>*fh298*</sup> mutants generated by Tilling (Draper et al. 2004). *nhsa*<sup>*fh298*</sup> carries a truncating

nonsense mutation at position Q217\* (C4524T) (Fig. 1B). Examination of embryos derived from an incross of *nhsa*<sup>fh298/+</sup> heterozygotes revealed no discernable morphological differences among siblings, suggesting that *nhsa* is dispensable for early development (<u>Fig. S1</u>).

We therefore decided to focus our efforts on zebrafish *nhsb* during embryogenesis. Using *in situ* hybridization, we found expression of *nhsb* in the eye during key stages of retinogenesis. Expression of *nhsb* was seen within proliferating retinal progenitor cells (RPCs) and in the lens at 24 hpf (Fig. 2A). By 48 hpf and 72 hpf, *nhsb* expression was downregulated in proliferating RPCs and not found in the ciliary marginal zone (CMZ). Instead, *nhsb* expression was detected in differentiated cells outside of the CMZ, with staining present in the retinal ganglion cell layer and inner plexiform layer of the retina (Fig. 2B-D).

We next examined a role for *nhsb* in zebrafish embryo development. To accomplish this, we designed splice-blocking antisense morpholino oligonucleotides (MO) to the exon2-intron2 boundary of the *nhsb* gene (*nhsb* MO1) (Fig. 1B). We found that *nhsb* MO1-injected embryos initially look indistinguishable from control embryos until 24 hpf, suggesting that knockdown of *nhsb* does not affect early development, eye field specification and/or separation. By 72 hpf, we observed a striking difference in the size and shape of the retina in *nhsb* morphants as compared to controls (Fig. 2E-H). *nhsb* morphants exhibited a small retina located on top of the lens (Fig. 2H). We did not observe any lens defects in *nhsb* MO1-injected embryos that were reminiscent of cataracts as observed in humans with Nance-Horan Syndrome (Burdon et al. 2003;

Brooks et al. 2004; Sharma et al. 2008; Coccia et al. 2009). We confirmed that injection of *nhsb* MO1 caused a mis-splicing of the *nhsb* transcript leading to retention of intron2 (Fig. S2). To ensure that the small eye phenotype was specific to knockdown of *nhsb*, we injected a second splice-blocking *nhsb* MO (*nhsb* MO2), targeted to the intron2-exon3 (I2E3) boundary (Fig. 1B). We found that *nhsb* MO2-injected embryos exhibited the same phenotype as that observed with *nhsb* MO1 (hereafter referred to as *nhsb MO*) (Fig. S3).

To further test whether the small eye phenotype was specific to disruptions in the *nhsb* gene, we used CRIPSR/Cas9 to generate insertion/deletion (indel) mutations at the *nhsb* locus. We designed a single-stranded guide RNA (gRNA) to target exon1 of the *nhsb* gene (Fig 1B). It was previously reported that injection of increasing amounts of gRNA can lead to bi-allelic disruptions in the targeted gene (Jao et al. 2013). We found that co-injection of *nhsb*<sup>ex1</sup> *gRNA* and *Cas9* mRNA into one-cell stage embryos led to growth defects in the eye of injected founder (F0) embryos (14%, *n* = 245), similar to that seen in *nhsb* morphants (Fig 2I,J). We chose five injected embryos displaying the small eye phenotype and validated indel mutations at the *nhsb* locus using a T7 endonuclease I (T7E1) assay; all assayed embryos showed high mutagenesis rates at the *nhsb* locus (Fig. S4). Taken together, our results demonstrate that the *nhsb* gene is expressed in the retina during key stages of retinogenesis, and that loss-of-function of *nhsb* leads to reduced retinal size in zebrafish.



#### Expression and function of NHS is conserved in Xenopus tropicalis

To test whether gene expression was conserved in other vertebrates, we examined a role for *nhs* in retina development in *Xenopus tropicalis*. We chose X. *tropicalis* because they are diploid and possess only one *nhs* gene. First, we made antisense *in situ* hybridization probes against the *Xenopus tropicalis nhs* gene (Fig. 1B).

Similar to zebrafish *nhsb*, the expression of *nhs* was detected in the retinal progenitor cells and lens of *Xenopus tropicalis* eyes at embryonic stage 31 (Fig. 3A). As retinogenesis proceeded, *nhs* expression in RPCs faded and was found concentrated in the plexiform layers of the retina (Fig. 3A-D).

To determine whether the role of Nhs in retina development was evolutionarily conserved among vertebrates, we designed splice-blocking MOs targeting the exon3intron3 boundary of *Xenopus tropicalis nhs* (*nhs* MO). We co-injected *nhs* MO along with rhodamine-dextran into one cell of a two-cell stage embryo. This experimental approach provided us with an internal control, since only half the embryo received *nhs* MO. We found that 96% of embryos (n = 56) injected with *nhs* MO had small retinas as compared to eyes on the contralateral side that did not receive morpholino (Fig. 3E-H). The efficacy of knockdown by the *nhs* MO in *Xenopus* was validated by RT-PCR (Fig. S5). These findings suggest that the role of NHS in retinal development is evolutionarily conserved since knockdown of *nhs* in frog and knockdown of *nhsb* in zebrafish results in a similar small eye phenotype. Taken together, the expression and function of Nhs during key stages of retinogenesis is conserved between *Xenopus tropicalis* and *Danio rerio*.



#### Nhs is required for proliferative growth of RPCs in the retina.

To further investigate the function of Nhs in retinal development we performed the rest of our analysis on zebrafish *nhsb* morphants. The earliest manifestation of the reduced eye size in *nhsb* morphants occurred between 24hpf - 48 hpf. This corresponds to the period of rapid RPC proliferation followed by neurogenesis and differentiation (He et al. 2012). To determine whether the reduced eye size was due to an increase in RPC cell death, we analyzed wild-type and *nhsb* morphant retinas for apoptotic cells. We evaluated cell death using an anti-cleaved caspase-3 antibody, a marker of apopotic cell death. We found virtually no dying cells in the retinas of either wild-type (0.187% (4/2137 cells, n = 6 embryos) or *nhsb*-deficient embryos (0.140%; 2/1427 cells, n = 6 embryos) at 36 hpf (Fig. 4A,B). Furthermore, *nhsb* morphant retinas did not display pyknotic morphology characteristic of dying cells (Fig. 4A,B). These results imply that the reduced eye size observed in *nhsb*-depleted embryos was not due to an increase in cell death.

To determine whether the small eye phenotype observed in *nhsb* morphants was due to a defect in RPC proliferation, we analyzed cryosections stained with phosphohistone 3 (PH3) antibody, a marker of M-phase of the cell cycle. At 24 hpf, a timepoint in which no morphological difference in eye size was observed, we found no significant difference in the proportion of PH3-positive cells in *nhsb* morphants compared to wild-type embryos (wt, 0.43 ± 0.05%, n = 538 cells, 4 embryos; *nhsb* MO, 0.51 ± 0.04%, n = 745 cells, 5 embryos)(Fig. 4B-I). At 48 hpf, however, the number of cells at M-phase was significantly fewer in the retina of *nhsb* morphants (0.92 ± 0.08%, n = 655 cells, 4 embryos) compared to wild-type (1.98 ± 0.18%, n = 2218 cells, 4 embryos) embryos (p < 0.05)(Fig. 4B-I). This suggests that reduced proliferation of RPCs may contribute to the lack of eye growth in the absence of *nhsb* function.



We next sought to determine whether differentiation of RPCs into neurons was affected by loss of *nhsb*. To accomplish this, we made use of *TgBAC(neurod1:EGFP)* transgenic zebrafish that expresses GFP under control of the promoter for *neuroD* (Thomas et al. 2012). This transgene is not expressed in proliferating RPCs, but is expressed in a subset of retinal neurons undergoing differentiation (Thomas et al. 2012). Some of the first differentiating neurons identified with this transgenic line can be seen around 34 hpf (Fig. 5A-C). Many GFP-positive neurons are present in the retina of wild-type embryos at this timepoint, and this number increased by 48 hpf (Fig. 4G-I). In contrast, by 34 hpf, no GFP-positive neurons are observed in *nhsb* morphants (Fig. 4D-F). By 48 hpf, however, a few GFP-expressing neurons can be visualized in *nhsb* morphant retinas (Fig. 4J-L), suggesting that the differentiation of neurons from RPCs is

not blocked but is delayed in *nhsb*-depleted embryos. These findings are consistent with the idea that *nhsb* knockdown mainly affects expansive growth of RPCs but does not inhibit neuronal differentiation.



Fig.5 nhsb knockdown leads to a delay in neuronal differentiation. (A) Assessment of GFP-positive cells in cryosections from TgBAC(neurod1:EG-FP) transgenic fish injected with nhsb MO at 34 hpf. At this timepoint, neurons are just differentiating in control embryos, whereas no GFP-positive cells were detected in nhsb morphant TgBAC(neurod1:EGFP) fish. (B) At 48 hpf, many neurons have differentiated in control embryos, whereas only a few neurons have terminally differentiated the nhsb morphants in TgBAC(neurod1:EGFP) fish.

## Loss of *nhsb* leads to a marked upregulation of $\Delta 113p53$ and *p21*, but not fulllength *p53*

To gain insight into the downstream function of Nhsb, we compared retinal expression profiles between wild-type and *nhsb* morphants. Total RNA was isolated from 40 zebrafish eyes dissected from wild-type and *nhsb* morphant embryos at 48 hpf for microarray analysis using Affymetrix zebrafish GeneChips. We used SAM analysis and found 1,212 significant genes that were differentially expressed in wild-type eyes compared to *nhsb* morphant eyes (Table S2A,B).

A subset of genes with significantly higher expression in *nhsb*-depleted eyes included genes that are markers of RPC identity such as *mz98/col15a1b* (*collagen typeXV, alpha 1b*), *ccnd1* (*cyclin D1*), *dlc* (*deltaC*), and *pold1* (*DNA polymerase delta 1*). (Fig. 6A). We also found a subset of genes whose expression was reduced in *nhsb* morphants compared to control, including markers of differentiated neurons such as *neuroD*, *opn1sw1* (*opsin1 short-wave-sensitive 1*), *rho* (*rhodopsin*), *calb2a* (*calbindin2a*), *snap25a* (*synaptosomal-associated protein, 25a*) (Fig. 6A). These findings are consistent with the idea that RPCs in *nhsb* morphant retinas proliferate slower, maintaining their identity as RPCs with delayed neuronal differentiation.

Surprisingly, functional annotation analysis using the Database for Annotation, Visualization and Integrated Discovery (DAVID) identified a number of genes that were differentially expressed between wild-type and *nhsb*-depleted retinas with known roles in the regulation of cell cycle progression, including the tumor-suppressor gene *tp53*  and its target gene *mdm2* (Fig. 6A). This suggests that loss of *nhsb* may regulate cell cycle progression through the p53 pathway.

To validate our microarray results, we performed semi-quantitative RT-PCR, using RNA extracted from dissected retinas of wild-type and *nhsb* morphants. We found that the levels of full-length *p53* transcripts were unchanged in *nhsb* morphants compared to wildtype controls, a result that differed from our microarray expression analysis (Fig. 6B). Whereas the microarray probe binds to the C-terminal *tp53* coding region, our RT-PCR primers amplify N-terminal *tp53* coding region from exon1 to exon4 (Table S1). However, using primers that specifically amplify the truncated  $\Delta 113p53$  transcript, we found a marked up-regulation of  $\Delta 113p53$  expression in *nhsb* morphant eyes compared to wild-type controls (Fig. 6B). Our results demonstrate that the elevated p53 transcripts detected by the Affymetrix probe, directed against the C-terminal coding region of p53, is due to the upregulation of the N-terminal truncated p53 variant  $\Delta 113p53$ , but not full-length *p53*.

We next asked whether other p53 target genes were affected by *nhsb* depletion. Semi-quantitative RT-PCR results showed that additional *p53* target genes were upregulated in *nhsb* morphant eyes compared to wild-type eyes, including *mdm2, ccng1 (cyclinG1), and cdkn1a/p21<sup>WAF1/CIP-1</sup>* (p21), a cyclin dependent kinase inhibitor that blocks the G1-to-S-phase transition (Cazzalini et al. 2010)(Fig. 5B). On the other hand, expression levels of the pro-apoptotic factor *bax* did not show apparent differences between *nhsb* morphants and control eyes (Fig. 5B). Taken together, these data are consistent with our histological data that the reduced eye size in *nhsb* morphants is not

caused by increased apoptosis but rather is due to a proliferation restriction imposed on

RPCs by elevated expression of the p53-target gene and cell-cycle checkpoint protein,

p21.



Knockdown of *p53*, reduces  $\Delta 113p53$  and *p21*, and rescues the *nhsb* morphant phenotype

Previous data from *Chen* and colleagues demonstrated that the induction of  $\Delta 113p53$  was p53-dependent (Chen et al. 2009). Our data suggest that *nhsb* may be a negative regulator of  $\Delta 113p53$  expression. Since  $\Delta 113p53$  is itself a transcriptional

target of full-length *p53* that binds to promoter elements in intron 4 (Chen et al. 2009), we reasoned that the small-eye phenotype seen in *nhsb* morphants may be rescued by knock-down of full-length p53. To determine whether full-length p53 is required for the elevated levels of  $\Delta 113p53$  caused by *nhsb* depletion, we co-injected *nhsb* MO2 with p53 MOATG that specifically targets the translation start site of the full-length p53 protein (Robu et al. 2007). We found that co-injection of *nhsb* MO and *p53* MO<sup>ATG</sup> rescued the small eye phenotype caused by injection of *nhsb* MO alone. (Fig. 7A). Splice blocking efficiency and knockdown of *nhsb* was confirmed by RT-PCR (Fig. S6) and phenotypes quantified (Fig. S7). Interestingly, co-injection of *nhsb* MO and *p53* MO<sup>ATG</sup> led to a reduced expression of  $\Delta 113p53$  and p21 transcripts, compared to *nhsb* MO alone (Fig. 6B). Our findings indicate that the upregulation of  $\Delta 113p53$  and p21 in nhsb-depleted retina is dependent on full length p53, consistent with the notion that  $\Delta 113p53$  and p21 are p53 target genes (Chen et al. 2009). Overall this result suggests that the proliferation restriction enforced on RPCs was alleviated when levels of  $\Delta 113p53$  and *p21* were reduced.

It was previously reported that  $\Delta 113p53$  antagonizes the apoptotic activities of p53 (Chen and Peng 2005; Chen et al. 2009), thereby promoting cell survival. To determine whether expression of  $\Delta 113p53$  variant is required for cell survival in *nhsb* morphants, we designed a second morpholino targeting the splicing site of the  $\Delta 113p53$  transcript at the exon ES - intron 4 splice junction ( $\Delta 113p53$  splMO). The activity of  $\Delta 113p53$  splMO is predicted to affect splicing of the  $\Delta 113p53$  transcript, but not full-length *p53* transcripts. Injection of either *p53* MO<sup>ATG</sup> or  $\Delta 113p53$  splMO alone had no

discernable effect on retinal development (data not shown). However, embryos coinjected with *nhsb* MO and  $\Delta 113p53$  splMO resulted in embryonic death, indicative of an increase in *p53*-mediated apoptosis when  $\Delta 113p53$  transcripts are absent.

Taken together, these results demonstrate that the upregulation of  $\Delta 113p53$  protects retinal progenitors from undergoing cell death in *nhsb* knockdown embryos, while imposing a proliferation restriction in a manner that likely involves other p53 target genes, such as the cell cycle inhibitor *p21*. Our data supports a model where *nhsb* is a negative regulator of  $\Delta 113p53$  expression and exerts its function in development though regulation of the p53 pathway.



#### Fig. 7 Knockdown of p53 rescues nhsb morphant phenotype and Δ 113p53 levels.

(A) Stereomicroscope images of zebrafish retina at 24 hpf and 48 hpf injected with nhsb MO alone or in combination with p53 MOATG (L = lens). Retinal size was not dramatically different in any condition at 24 hpf. However, injection of p53 MOATG targeting full-length p53 transcripts rescued retinal size in nhsb morphants. (B) Semi-quantitative **RT-PCR** examination of p53 target genes. Note that injection of p53 MOATG inhibited the upregulation of ∆113p53 and p21 transcripts caused by nhsb-depletion. (C) Injection of Δ113p53 MO that specifically targets the ∆113p53 variant and not full-length p53 led to elevated cell death.

#### Discussion

The vertebrate retina develops from a small pool of retinal progenitor cells (RPCs) comprising the optic cup. Just prior to the onset of retinal neurogenesis, RPCs undergo cycles of rapid self-replication to expand the tissue (Agathocleous and Harris 2009; He et al. 2012). The proliferative capacity of RPC's are diminished when key transcription factors like Rx1, Pax6, Chx10, and Sox2 are inhibited, whereas activation of these transcription factors often lads to increased mitosis and a larger retina (Marguardt et al. 2001; Green et al. 2003; Philips et al. 2005; Van Raay et al. 2005; Taranova et al. 2006; Zaghloul and Moody 2007; Oron-Karni et al. 2008). Temporal expression of bHLH proteins like neurogenin 2 (Ngn2), atonal homolog 5 (ATH5) and NeuroD trigger cell cycle exit to initiate specification and differentiation in a highly coordinated manner (Agathocleous and Harris 2009). Extrinsic signals from the retina microenvironment also affects division and differentiation of RPCs. Ligands like Sonic Hedgehog (Shh) promotes rapid proliferation of RPCs by shortening cell cycle length through the activation of G1 and G2 phase cyclins (Wang et al. 2005; Locker et al. 2006; Agathocleous et al. 2007). It has been proposed that Shh stimulates the transition of early RPC from slow cycling to faster cycling in advance of terminal differentiation (Locker et al. 2006; Agathocleous et al. 2007). Our results demonstrate that knockdown of Nhs in zebrafish and Xenopus leads to a reduction in proliferation of RPCs before the onset of differentiation. We propose that loss of Nhs function leads to an increase in cell cycle length that may be due to the elevated levels of cyclin-dependent kinase inhibitors, such as p21, as a result of p53 pathway activation. Our results show that Nhs-depletion does not inhibit, but rather delays the differentiation of RPCs. Although differentiation proceeds, it remains unclear whether loss of Nhs function affects the differentiation of RPCs into all possible cell types in the retina. Nevertheless, our findings indicate Nhs regulates overall retinal size by promoting the faster cycling of RPCs associated with the expansion of the RPC population in advance of terminal differentiation.

Although it is clear that the p53 tumor suppressor plays important roles in cancer progression, a role for p53 (or any or the p53 isoforms) in embryonic development or development of the nervous system is less clear. p53 acts primarily as a transcription factor, although it has been shown to have a cytoplasmic roles (Vousden 2006; Endo et al. 2008). In response to DNA damage or other cellular stresses, there are two main outcomes of p53 activation, namely cell cycle arrest or apoptosis. In the nervous system, p53 plays a key role, along with other p53 family members, p63 and p73, in regulating survival and apoptosis in the developing and adult nervous system (Miller et al. 2000; Jacobs et al. 2004). p53 becomes a major pro-apoptotic proteins eliciting naturally occurring cell death in peripheral neurons (Aloyz et al. 1998; Jacobs et al. 2005) and neural progenitor cells (Dugani et al. 2009; Fatt et al. 2014). Consistent with this, mice deficient in DNA ligase 4 (Lig4) or XRCC4, two components of the nonhomologous end-joining (NHEJ) pathway that repairs double-stranded DNA breaks, exhibit neuronal apoptosis that has been shown to be p53-dependent (Frank et al. 1998; Gao et al. 1998). A screen of zebrafish mutants harboring mutations various categories of cell essential genes, such as DNA replication, transcription, telomere

maintenance, ribosome biogenesis, splicing, chaperoning, endocytosis, and cellular transport, all share an upregulation of p53 and excessive neural apoptosis (Danilova et al. 2010). Activation of the p53 pathway then is likely responsible for a wide range of defects seen in human congenital disorders (Mendrysa et al. 2011).

Few studies, however, have investigated the role of p53 isoforms, such as  $\Delta$ 113p53/ $\Delta$ 113p53, in embryonic development. The zebrafish protein  $\Delta$ 113p53, and its human counterpart,  $\Delta$ 133p53, are N-terminally truncated forms of p53 that are missing the transactivation domain, and MDM2-interacting motif, with a partial deletion of the DNA-binding domain (Bourdon et al. 2005; Chen et al. 2005, 2009). Transcriptional expression of  $\Delta$ 113p53/ $\Delta$ 133p53 is initiated by an alternative p53 promoter in intron 4 that is completely dependent on full length p53 protein (Bourdon et al. 2005; Chen et al. 2009; Aoubala et al. 2011). Δ113p53/Δ133p53 is strongly induced by DNA double strand breaks, where it antagonizes p53-mediated apoptosis(Gong et al. 2015). Previous work had shown that  $\Delta$ 113p53 does not act in a dominant-negative fashion, but by differentially modulating p53 target gene activation to protect cells from apoptosis (Chen et al. 2009; Ou et al. 2014). In zebrafish,  $\Delta 113p53$  was identified in the analysis of a zebrafish mutant def, a pan-endodermal-enriched factor that is essential for the expansive growth of digestive organs (Chen et al. 2005). In *def* null mutants,  $\Delta 113p53$ transcripts were upregulated and induced the expression of p53-responsive genes to trigger arrest of cell cycle, but not apoptosis, leading to reduced organ growth (Chen et al. 2005, 2009). We identified a similar role for *nhsb* in expansion growth of the retina. In the absence of *nhsb*,  $\Delta 113p53$  expression increased, which concomitant increase of the

p53 target gene *p21*. Increased levels of *p21* enforced a proliferation restriction on RPCs, while the pro-apoptotic gene *bax* was not increased in hypoplastic retinas. Our findings are in agreement with the previously reported role for  $\Delta 113p53$  as regulator of *p21*, but not *bax* (Chen et al. 2005, 2009). These results suggest that i) Nhs is required in retinal progenitor cells to keep levels of  $\Delta 113p53$  low, and that ii) loss of *nhs* causes a small retina due to cell cycle arrest, but not apoptosis. Indeed, when  $\Delta 113p53$  were depleted, loss of *nhsb* led to an increase in cell death.

How does Nhs regulate the  $\Delta 113p53$  transcript levels? Our data suggest that Nhs is a negative regulator of  $\Delta 113p53$  expression. Nhs could regulate  $\Delta 113p53$  levels directly or indirectly through modulation of the p53 pathway. Our experiments indicate full length p53 is required for the micropthalmia and  $\Delta 113p53$  induction caused by *nhsb* deficiency. However, we show that transcript levels of full-length *p53* are unchanged in *nhsb* morphants, suggesting that Nhs does not regulate p53 at the level of transcription. One possibility is that Nhs regulates the steady state levels of p53 protein, and in the absence of Nhs, p53 levels would accumulate and elevate expression of  $\Delta 113p53$  and *p21*. Alternatively, Nhs could limit the transcriptional activity of p53 without affecting protein levels. It was recently reported that Def is a nucleolar protein that mediates p53 protein degradation through a proteasome-independent pathway (Tao et al. 2013). Future studies will be aimed at uncovering the mechanism of how Nhs modulates the p53 pathway.

Finally, our study uncovers a developmental role for Nhs that provides an underlying mechanism for pathology associated with Nance-Horan Syndrome (NHS).

Affected males have severe bilateral congenital dense nuclear cataracts that lead to profound vision loss and usually require surgery at an early age (Burdon et al. 2003; Huang et al. 2007; Sharma et al. 2008; Coccia et al. 2009). Despite cataract surgery during infancy, many NHS patients continue to exhibit visual impairment that may be caused by an underlying retinopathy (Mathys et al. 2007; Ding et al. 2009). Our findings that Nhs is essential for proper retinal development may clarify why NHS patients continue to exhibit ocular abnormalities and vision loss, even when cataracts have been corrected by surgery. NHS is caused predominantly by truncating (nonsense) mutations in the NHS gene that have been found in each of the 8 exons. There is no apparent clinical difference between mutations in C-terminal exons compared to N-terminal exons (Tug et al. 2013). Previous reports have suggested that humans have at least three isoforms of NHS transcripts generated by the use of alternative transcriptional start sites, arising from exon1, exon1a, and exon1b (see Fig 1B). Transcripts encoded from exon1 is known to be important in the pathogenesis of NHS, because patient nonsense mutations identified in exon1 are only predicted to affect this isoform (Burdon et al. 2003; Brooks et al. 2004; Huang et al. 2006; Hong et al. 2014). This exon1 is evolutionarily conserved in vertebrates. Our results presented here using CRISPR genome editing with guide RNAs directed against exon1 of the zebrafish nhsb gene showed a similar microphalmia compared with morpholino oligonucleotide-mediated knockdown. These results are consistent with the human molecular genetics demonstrating that NHS isoforms encoded from exon1 are relevant to the pathogenesis of Nance-Horan Syndrome.

## **Materials and Methods**

#### Zebrafish and Xenopus tropicalis Husbandry

Zebrafish (Danio rerio) were maintained according to standard procedures and staged as previously described (Kimmel et al. 1995). The nhsa<sup>th298</sup> mutant line carries a truncating nonsense mutation at position Q217\* and was generated by TILLING (Draper et al. 2004). The neurod:EGFP transgenic line is registered as *TgBAC(neurod1:EGFP)nl1Tg* at The Zebrafish International Resource Center (ZIRC)(Obholzer et al. 2008). Xenopus tropicalis embryos were obtained and cultured using standard methods (Sive et al. 2000).

#### Phlyogenetic analysis

Coding DNA sequences were gathered from, Ensembl (http://www.ensembl.org). Splice variants were checked by Mouse Genome Informatics (http://www.informatics.jax.org/), Zfin (http://zfin.org/), Xenbase (http://www.xenbase.org) and FlyBase (http://flybase.org/). *Xenopus* and zebrafish genes were manually curated using UCSC genome browser (https://genome.ucsc.edu/), Xenbase and Zfin. The following genes were used in our analysis:

(ENSG00000135540), NHSL2 human: NHS (ENSG00000188158), NHSL1 (ENSG00000204131). Mus musculus: NHS (ENSMUSG00000059493), NHSL1 (ENSMUSG00000039835), NHSL2 (ENSMUSG00000079481). Xenopus tropicalis: NHS (ENSXETG0000001112), NHSL1 (ENSXETG0000006237), NHSL2 (ENSXETG00000022580). Danio rerio: nhsa (ENSDARG00000070227), nhsb (ENSDARG00000079977), nhsl1a (ENSDARG00000054537), nhsl1b (ENSDARG00000042627), nhsl2a (ENSDARG00000089066), nhsl2b (ENSDARG00000090164). <u>Drosophila melanogaster</u>: gukh (FBgn0026239). Above genes were entered into ClustIW2 Phylogeny (http://www.ebi.ac.uk/) and aligned by multiple sequence alignment. A Newick output was generated using the Neighbourjoining clustering method. Cladogram of Coding DNA was composed using Genious 7.0.6 software.

#### Whole-mount in situ hybridization

RNA in situ hybridization was carried out as previously described (Walsh et al., 2011). The gene specific probe fragments to the 3'UTR region of *nhs* genes were obtained by PCR (see Table S1 for primer pairs) from cDNA of 24 hpf zebrafish wild type embryos or early-stage *X. tropicalis* embryos. PCR products were cloned into pCR4-Blunt, sequenced. Sense and antisense riboprobes were transcribed from linearized plasmids and labeled with digoxigenin (DIG RNA-labelling reagents, Roche Biochemicals). Primers used to amplify zebrafish *nhsb* were 5' tgatctaccttttctcccatgccatt 3' and 5' tctcacacacacagaggctcca 3'. Primers used to amplify *Xenopus tropicalis nhs* were 5' tgaccagccacattgattatag 3' and 5' ccttaaaaagcaggccacagtt 3'.

#### **Morpholino injections**

Antisense MOs were injected at the 1-cell stage in zebrafish embryos and into one cell of a two-cell stage *Xenopus* embryo. Microinjections of antisense MOs were carried out

using an Eppendorf microinjector and Zeiss stereoscope, as described (Dickinson and Sive, 2009). Morpholinos were supplied by Gene Tools and were as follows:

*zebrafish nhsb*: MO1 E2I2 5'CCAGAAGACCCATCAGTACCTCTGT3', 4 ng, MO2 I2E3 5' CTGCTGCTGGTCTGTGAAGCAAACA 3', 3.5 ng, *Xenopus tropicalis nhs:* MO E3I3 5' GTGTGTAATATATCTTACCTCTCCG 3', 8.5 ng. The *p53* MO<sup>ATG</sup> targets the translation start site of full-length p53 as described (Robu et al. 2007): 5'GCGCCATTGCTTTGCAAGAATTG 3', 3 ng. A morpholino that targets the splice junction from exon ES to intron 4 of p53 and disrupts splicing of  $\Delta$ 113p53 isoform while leaving full-length p53 intact: spIMO 5' TGTCTTTCAAATGTCTTACCCTCC 3', 3 ng.

#### **CRISPR** Design and injection

We designed guide RNA (gRNA) sequence directed against exon1 of *nhsb* (5' GGTCCGGGATAGAGCCACAT 3'). For making gRNA, the guide sequence template was generated by annealing and filling-in two oligonucleotides: guide oligo (5' aattaatacgactcactataGGTCCGGGATAGAGCCACATgttttagagctagaaatagc3') and scaffold template

(5'GATCCGCACCGACTCGGTGCCACTTTTTCAAGTTGATAACGGAC

TAGCCTTATTTTAACTTGCTATTTCTAGCTCTAAAAC3'). The resulting gRNA template that contains a T7 promoter, was purified using DNA clean columns (ZymoResearch). *gRNA* was generated by in vitro transcription using T7 RNA polymerase (Megascript T7 Kit, Invitrogen). After transcription, gRNA was column purified (ZymoResearch). For making nls-zCas9-nls RNA, the template DNA (pT3TS-

nls-zCas9-nls; kind gift of Wenbiao Chen (Addgene plasmid # 46757) was linearized by XbaI, phenol-chloroform extracted followed by ethanol precipitation. Capped nls-zCas9nls mRNA was synthesized using mMessage T3 kit (Invitrogen) and purified using Illustra Probe-quant columns (GE Healthcare). The mix of gRNA (100ng/ $\mu$ L) and nlszCas9-nls mRNA (300 ng/ $\mu$ L) was injected into one-cell stage embryos.

#### **T7 Endonuclease I Assay**

Genomic DNA was prepared from whole embryos at 24 hpf. A short stretch of genomic region (~735 bp) flanking the target site was PCR amplified from genomic DNA. PCR product was denatured and slowly allowed to re-anneal to encourage heteroduplex formation. The re-annealing procedure consisted of a 5 minute denaturation step, followed by cooling to 85°C at -2°C per second and further to 25°C at a rate of -0.1°C per second. The re-annealed amplicon was then digested with 20 units of T7 endonuclease I (New England Biolabs) at 37°C for 15 minutes. The reaction was stopped by adding 1  $\mu$ L of 0.5M EDTA. The sample was run through a 2% agarose gel and visualized by ethidium bromide staining.

#### **Reverse Transcription-Polymerase Chain Reaction (RT-PCR)**

Splice-blocking MOs were validated by RT-PCR. Primers used to validate are listed in Supplementary Table 1. Briefly, total RNA was isolated from whole embryos using an RNeasy kit (Qiagen) and genomic contamination was eliminated by on-column DNA digestion (Qiagen). First Strand cDNA was synthesized using the Protoscript M-MuLV First-Strand cDNA Synthesis Kit (New England Biolabs). Equal concentrations of RNA (1ug) from all developmental timepoints were used in the cDNA synthesis reaction. Concentration of cDNA across developmental timepoints was validated using primers specific to the ornithine decarboxylase (ODC) gene or *elf1a*. RNA extracted from retinas were used to validate array data. For semiquantitative RT-PCR, the amount of template and the number of PCR cycles were optimized to ensure that the reactions were in the linear range of amplification. The primer pairs and detailed PCR conditions used to amplify each of these genes are listed in Table S1.

#### Affymetrix Array

We dissected 40 eyes each from wildtype or *nhsb* MO2-injected embryos at 48hpf. This experiment was performed in duplicate. Dissections were done in Half Ringers Solution (2.2mM CaCl2, 5.6mM KCl, 154 mM NaCl, 2.4mM NaHCO3, 2 mM Tris-Cl, adjust to pH 7.4) on ice. Total RNA was extracted from pooled tissues using RNeasy kit (Qiagen) and treated with DNAse I. We used 200ng input total RNA for cRNA amplification using the Affymetrix One Cycle kit. Amplicons were fragmented, hybridized to GeneChip Zebrafish Genome Arrays and scanned (Affymetrix). Quality control of total RNA, amplified RNA, and fragmented RNA was checked using Experion (Bio-Rad). GeneChip arrays were scanned on an Affymetrix Scanner 3000 7G scanner. Data was analyzed by Affymetrix Expression Console software. Raw intensity data was exported and filtered with a lower cut off value of 2 to assure that probe intensities data were generated within the dynamic range of fluorescence. Data was normalized by Robust

Multi-array Average (RMA) analysis to account for a possible probe effect, and uploaded into TMEV for Significance Analysis of Microarrays (SAM) analysis. The resulting 1,212 significant genes were entered into DAVID for further gene ontology analysis.

#### Immunostaining and Imaging

Embryos were fixed in 4% PFA overnight at 4°C, incubated in 30% sucrose and saturated in OCT (Tissue-Tec), frozen on cryostat stages prior to storage at -20°C. Coronal sections of 10uM were cut on a cryostat and mounted to Colorfrost Plus, microscope slides (Fisher). Immunostaining was performed on sections with the following antibodies: anti-phosphohistone 3 (pH3) (1:1000 Cell Signaling); anti-GFP (1:2000, Torrey Pines); cleaved caspase 3 (1:500) (Cell Signaling). Primary antibodies were detected using secondary antibodies: goat anti-mouse Alexa 564 or goat anti-rabbit Alexa 488 (Life Technologies). Nuclei were labeled with Dapi. Sections were mounted in 100% glycerol and coverslipped. Confocal images of labeled cryostat sections or whole-mount embryos were obtained on a Carl Zeiss Spinning Disk Laser Confocal Observer Z1 (VCU) using either 20X or 63X objectives.

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**Fig. 1 Phlyogeny and genomic architecture of NHS family genes** (A) Phylogenetic tree for the nhs family of genes including nhs, nhs-like 1 (nhsl1), and nhs-like 2 (nhsl2) and their common homolog in Drosophila, guanylate kinase holder (gukh). Full protein coding sequences were used for each family member. The initial letter hs denotes human, mm mouse, xt Xenopus tropicalis, dm Drosophila melano-gaster, and dr Danio rerio. (B) Conserved genomic structure of nhs genes in vertebrates showing relative positions of exons and alternative translation start sites. Dark boxes represent exons, open boxes represent untranslated regions (UTRs), star represents location of nonsense mutation (fh298) in nhsa, red boxes represent predicted morpholino oligonucleotide binding region, yellow dot represents nhsb guide RNA sequence target. (C) Temporal expression patterns of nhsa and nhsb in zebrafish embryos. Polymerase chain reaction from zebrafish cDNA at various timepoints using primers designed to nhsa, nhsb, and the control ornithine decarboxylase (odc).



## Control

# nhsb MO1







## Control

## CRISPR (nhs exon1)



**Fig. 2** Loss of nhsb function leads to a reduced eye size in zebrafish. Whole-mount in situ hybridization using a nhsb riboprobe to examine the expression patterns nhsb at 24hpf (A), 48 hpf (B), 72 hpf (C), 96 hpf (D). Stereomicroscope photomicrograph of embryos at 72 hpf from control (E) and nhsb morphants (F). In comparison to control embryos (E,G), embryos injected with nhsb MO1 display a smaller eye size. Stereomicroscope photomicrographs of embryos at 48 hpf from control (I) and F0 embryos injected with Cas9/gRNA targeted to exon1 of nhsb (J).



# Control

# nhsb MO





## Fig. 3 Loss of nhs function causes a small eye in frog. Vibratome

sections of Xenopus tropicalis embryos stained by whole-mount in situ hybridization using a probe against nhs at Stage 31 (A), Stage 33 (B), Stage 37 (C), and Stage 39 (D). Stereomicroscope photomicrographs of Xenopus tropicalis embryos at stage 38 of control (E,G) and nhs MO-injected embryos (F,H). Note the small eye phenotype in nhs morphant frogs.



# Fig. 4 depleted. (\*\* p < 0.05, Student's t test)

Reduced proliferation in retinal progenitor cells but not cell death following nhsb depletion. Cryosections of zebrafish retina stained for Dapi (blue) and anti-cleaved caspase-3 (CC3) (green) in control wild-type (A), and nhsb morphants (B). Despite having less cells, no increase in CC3 staining is evident in nhsb-depleted embryos compared to control. (C-H) Cryosections of zebrafish retina stained for Dapi (blue) and anti-phosphohistone3 (PH3) (red) at 48 hpf. (I) Quantitation of PH3-positive cells in the retina of control and nhsb morphants. The number of PH3-positive cells are significantly reduced in nhsb-morphant retinas compared to control wild-type retinas at 48 hpf indicating a decrease in proliferating cells when nhsb is









## Fig. 5

nhsb knockdown leads to a delay in neuronal differentiation. (A) Assessment of GFP-positive cells in cryosections from TgBAC(neurod1:EGFP) transgenic fish injected with nhsb MO at 34 hpf. At this timepoint, neurons are just differentiating in control embryos, whereas no GFP-positive

## cells were detected in nhsb morphant TgBAC(neurod1:EGFP) fish. (B) At 48 hpf, many neurons have differentiated in control embryos, whereas only a few neurons have terminally differentiated in the nhsb morphants TgBAC(neurod1:EGFP) fish.

A					
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	de			cacnb1	
	L			snap25a	
	or			rc3	
	lei			opn1sw1	



**Fig. 6** Δ113p53 but not full length p53 is upregulated in nhsb morphants. (A) Clustering and heat map of gene transcripts regulated by

nhsb-depletion identified in microarray experiments. Red and green shading, respectively, indicates the highest and lowest expression even by peer review) is the author/funder, who has granted bioRxiv a liperse to display the preprint in perpirituity. It is made available under even by peer review) is the author/funder, who has granted bioRxiv a liperse to display the preprint in perpirituity. It is made available under even by a co-By NC-By of the figure. Following knockdown of nhsb, markers of RPC identity were elevated, whereas marker of terminally differentiating neurons were downregulated, and genes associated with the p53 pathway were epregulated compared to wild-type control retinas (B) Semi-quantitative RT-PCR examination of expression of various p53 response genes in nhsb morphant retina and wild-type control retinas. nhsb morphants had elevated levels of  $\Delta 113p53$ , p21, mdm2, and cyclin G1, but full-length p53 and bax remained unchanged



Fig. 6 Δ113p53 but not full length p53 is upregulated in nhsb morphants. (A) Clustering and heat map of gene transcripts regulated by nhsb-depletion identified in microarray experiments. Red and green shading, respectively, indicates the highest and lowest expression levels, as indicated in the scale bar at the bottom of the figure. Following knockdown of nhsb, markers of RPC identity were elevated, whereas marker of terminally differentiating neurons were downregulated, and genes associated with the p53 pathway were epregulated compared to wild-type control retinas (B) Semi-quantitative RT-PCR examination of expression of various p53 response genes in nhsb morphant retina and wild-type control retinas. nhsb morphants had elevated levels of  $\Delta 113p53$ , p21, mdm2, and cyclin G1, but full-length p53 and bax remained unchanged