SOX2 is a dose-dependent regulator of retinal neural progenitor competence

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Approximately 10% of humans with anophthalmia (absent eye) or severe microphthalmia (small eye) show haploid insufficiency due to mutations in SOX2, a SOX-B1-HMG box transcription factor. However, at present, the molecular or cellular mechanisms responsible for these conditions are poorly understood. Here, we directly assessed the requirement for SOX2 during eye development by generating a gene-dosage allelic series of Sox2 mutations in the mouse. The Sox2 mutant mice display a range of eye phenotypes consistent with human syndromes and the severity of these phenotypes directly relates to the levels of SOX2 expression found in progenitor cells of the neural retina. Retinal progenitor cells with conditionally ablated Sox2 lose competence to both proliferate and terminally differentiate. In contrast, in Sox2 hypomorphic/null mice, a reduction of SOX2 expression to <40% of normal causes variable microphthalmia as a result of aberrant neural progenitor differentiation. Furthermore, we provide genetic and molecular evidence that SOX2 activity, in a concentration-dependent manner, plays a key role in the regulation of the NOTCH1 signaling pathway in retinal progenitor cells. Collectively, these results show that precise regulation of SOX2 dosage is critical for temporal and spatial regulation of retinal progenitor cell differentiation and provide a cellular and molecular model for understanding how hypomorphic levels of SOX2 cause retinal defects in humans.

[Keywords: SOX2; allelic series; retinal progenitor identity; dosage regulation; anophthalmia, microphthalmia]

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Seven main classes of retinal cell types (rod, cone, bipolar, horizontal, amacrine, ganglion, and Müller glial cells) are produced from a pool of multipotent retinal progenitor cells (RPC). The overall size of the retina, and the proportion of each of these cell types contained therein is essential for proper visual processing. To ensure that the adult retina forms appropriately during development, RPCs transition through states of developmental competence by coordinating cell cycle exit and cell fate specification to generate an ordered array of uniquely fated cell populations [Dyer and Cepko 2001; Fujita 2003; Pearson and Doe 2004; Zhang et al. 2004; Kageyama et al. 2005]. When these processes are disrupted, as observed in some cases of human microphthalmia and anophthalmia [Zigman and Paxhia 1988; Loosli et al. 1998; Fantes et al. 2003; Driver et al. 2005; Fitzpatrick and van Heyningen 2005; Labadie et al. 2005; O’Brien et al. 2005; Ragge et al. 2005a,b; Xu et al. 2005], vision is severely compromised. Identifying the molecular mechanisms that allow some progenitors to continually divide yet promote neighboring cells to stop dividing and differentiate at appropriate times during development is pivotal to the understanding of both normal and defective retinal development.

It has recently been shown that mutations in SOX2 a SOXB1-HMG box transcription factor whose expression universally marks neural stem and progenitor cells throughout the CNS including the neural retina [Collignon et al. 1996; Zappone et al. 2000; D’Amour and Gage 2003; Ellis et al. 2004; Ferri et al. 2004], are associated with retinal and ocular malformations in humans. The resulting haploid insufficiency at the SOX2 locus occurs in ~10% of human individuals with anophthalmia or severe microphthalmia [Fantes et al. 2003; Fitzpatrick and van Heyningen 2005; Hagstrom et al. 2005; Ragge et al. 2005a,b; Zenteno et al. 2005]. Most mutations identified to date are point mutations leading to truncations of SOX2, while a smaller class of mutations includes microdeletions and missense point mutations. Interestingly, all mutations produce hypomorphic conditions, where residual SOX2 expression and function are still preserved, albeit at lower levels, leading to the highly variable severity of the clinical phenotype. In this regard, the SOX2 mutations in humans and the clinical consequence of reduced functional levels of SOX2 suggest a dosage-dependent role for SOX2 during retinal progenitor differentiation.

To date, the importance of SOX2 in the nervous system has been highlighted by misexpression and domi-
nant interfering studies in mouse cell lines, *Xenopus*,
and chick embryos, which suggests that SOX2 maintains
neural progenitor identity [Mizuseki et al. 1998; Kishi
et al. 2000; Bylund et al. 2003; Graham et al. 2003; Pevny
and Placzek 2005; Van Raay et al. 2005]. However,
the lethality of Sox2-null mutant mice at the preimplanta-
tion stage [Avilion et al. 2003] combined with the over-
ap in expression and potential functional redundancy
of the three highly related SOX1 factors [SOX1, SOX2,
and SOX3] in CNS progenitors [Collignon et al. 1996; Pevny
et al. 1998; Weiss et al. 2003; Rizzoti et al. 2004; Tanaka
et al. 2004] have precluded genetic evaluation of the spe-
cific role of SOX2 in neural progenitor cells.

Unlike the extensive overlap of their expression in the
rest of the CNS, SOX1 factors display unique patterns
of expression in the developing eye [Kamachi et al. 1998,
2001; Nishiguchi et al. 1998; Uchikawa et al. 1999, 2003;
Le et al. 2002]. SOX1, SOX2, and SOX3 are initially ex-
pressed in the anterior neural plate and invading optic
vesicle. However, during the formation of the optic
cup, SOX1 and SOX3 are down-regulated, while SOX2 is
maintained and restricted to neural retinal cells [Kama-
chi et al. 1999, 2001; Le et al. 2002]. Thus, the selective
expression of SOX2, and not SOX1 or SOX3, in the neu-
ral retina, coupled with cellular and molecular parallels
between ventricular zone and retinal CNS progenitor dif-
ferentiation, provides an ideal in vivo model system to
evaluate the unique contribution of SOX2 in neural pro-
genitor populations [Collignon et al. 1996; Kamachi et al.
1998; Nishiguchi et al. 1998; Le et al. 2002; Uchikawa et
al. 2003; Blackshaw et al. 2004].

To dissect out the retinal-specific roles of SOX2 and to
determine the effect of SOX2 dosage on these roles, here
we report the examination of systematic dosage reduc-
ton of SOX2 on retinal progenitor differentiation using
conditional/null, compound hypomorphic/null, and het-
erozygous/null and wild-type mice. First, we show that
conditional ablation of SOX2 in the mouse retina causes
the complete loss of neural progenitor competence to
divide and differentiate. Second, and in contrast with
complete ablation of SOX2, hypomorphic levels of SOX2
expression trigger aberrant retinal progenitor differentia-
tion, resulting in variable microphthalmic phenotypes.
In addition, using two independent approaches we dem-
strate that SOX2, acting as a transcription factor, me-
diates its effects by directly regulating the expression
levels of NOTCH1. These studies provide direct cellular
and molecular evidence that alterations in the levels of
SOX2 regulate the choice between maintenance of pro-
genitor cell identity and differentiation. Moreover, these
data reveal a molecular basis for retinal defects in hu-
mans, which present as anophthalmia and microphthal-
mia syndromes, as a consequence of hypomorphic levels
of SOX2.

Results

**SOX2 defines retinal progenitor cells**

Previous studies have shown that SOX2 expression
marks proliferating neural progenitors and is down-regu-
lated in the CNS ventricular zone concomitant with
their differentiation [Ellis et al. 2004; Ferri et al. 2004;
Rizzoti et al. 2004; Tanaka et al. 2004]. We therefore
addressed whether SOX2 expression is regulated in a
similar dynamic manner in RPCs. RPC differentiation
results in the formation of discrete layers; progenitor
cells are retained in the outer retinoblast layer (RBL) [ar-
row in Fig. 1B], while cells that exit the cell cycle are
located in the inner ganglion cell layer (GCL) [arrow in
Fig. 1D]. Upon exiting the cell cycle, retinal progenitors
differentiate into six defined neuronal cell types in a pre-
cise temporal and spatial manner [Cepko et al. 1996, Reh
and Fischer 2001]. Retinal ganglion cells (RGCs) are
generated first, followed by overlapping phases of develop-
ment for horizontal cells, cones, amacrine cells, rods,
and bipolar cells. Using a Sox2EGFP-reporter mouse line
in which the Sox2 coding region is replaced by an-
enhanced green fluorescent protein [EGFP] such that EGFP
fluorescence recapitulates SOX2 expression [Ellis et al.
2004] we show that, as in the CNS ventricular zone,
SOX2 and EGFP [Fig. 1A,E] expression in the embryonic
retina coincides with markers of cell proliferation such as
Proliferating Cell Nuclear Antigen [PCNA] [Fig. 1B,F].
However, in contrast to CNS ventricular zone progeni-
tors that also express SOX1 and SOX3, retinal neural pro-
genitors exclusively express SOX2 [Fig. 1C,G]. Coinc-
dent with retinal cell differentiation, SOX2 (and EGFP)
expression is down-regulated such that its expression is
mutually exclusive of the general neuronal marker neu-
ron-specific class III βTUBULIN [βTUBULIN III] in the
GCL [Fig. 1D,H]. Subsequently, SOX2 (and EGFP)
expression is not detected in cells expressing
specific markers associated with each of the mature reti-
nal cell types including NEUROFILAMENT, a marker of
RGCs and horizontal cells [Fig. 1I,L], RHODOPSIN, a
marker of rod cells [Fig. 1J,M], and Protein Kinase C α
[PKC], a marker of bipolar cells [Fig. 1K,N]. Consistent
with the expression pattern of Sox2 in the chick [Le et al.
2002], SOX2 is maintained in a small subset of cells iden-
tified as displaced amacrine by the coexpression of
ISLET1 [Fig. 1O,R] and CALRETININ [Fig. 1P,S]. In con-
trast to its down-regulation in post-mitotic neuronal
cells, SOX2 is maintained in Müller glia, a nonneuronal
cell type of the mature retina, marked by Cellular Reti-
nal-Binding Protein [CRALBP] [Fig. 1Q,T] [Eisenfeld et
al. 1985] and Glial Fibrillary Acidic Protein [GFAP] [data
not shown] expression.

These analyses demonstrate that SOX2 expression in
both the neural retina and ventricular zone of the CNS is
inversely correlated with the progression of neuronal dif-
fferentiation, suggesting that there is a shared mecha-
nism regulated by SOX2.

*A dosage-dependent role for SOX2 during retinal
progenitor differentiation*

To analyze the effects of decreasing levels of SOX2 in
neural progenitor cells, we generated an allelic series
of Sox2 mutations in the mouse including a null
(Sox2EGFP), a conditional null (Sox2COND), and two hy-
pomorphic (Sox2<sup>LP</sup> and Sox2<sup>IR</sup>) alleles. Sox2<sup>EGFP</sup> is a null allele in which the Sox2 ORF has been substituted with an EGFP expression cassette [Ellis et al. 2004]. The Sox2<sup>COND</sup> allele contains the Sox2 ORF flanked by lox<sup>P</sup> sites such that CRE-mediated recombination results in removal of the entire SOX2 coding sequence and generates a null allele (Sox2<sup>ACOND</sup>) [Fig. 2A,B], and Sox2<sup>LP</sup> and Sox2<sup>IR</sup> were constructed by the insertion of PGK–neo–loxP and IRESdsRED-loxP-flanked-PGKneo expression cassettes, respectively, 3′ of the Sox2 ORF [Fig. 2A,C,D].
Sox2<sup>LP</sup> and Sox2<sup>IR</sup> alleles effectively act as hypomorphic alleles, displaying <40% activity in a Sox2-null genetic background. Sox2 mRNA (data not shown) and protein levels in cells isolated from embryonic day 14 (E14) CNS cortex (Fig. 3A, B) and eyes (Fig. 3C, D) are decreased to 15%–30% and 20%–40% of wild type in Sox2<sup>EGFP/LP</sup> and Sox2<sup>EGFP/IR</sup> mice, respectively.

Animals heterozygous for Sox2<sup>+/EGFP</sup>, Sox2<sup>+/COND</sup>, Sox2<sup>+/IR</sup>, and Sox2<sup>+/LP</sup> and homozygous for Sox2<sup>COND/COND</sup>, Sox2<sup>IR/IR</sup>, and Sox2<sup>LP/LP</sup> are viable, born in appropriate Mendelian ratios, and are phenotypically and morphologically indistinguishable from wild-type mice (Fig. 2E; Supplementary Fig. 1). However, Sox2 hypomorphic/null compound heterozygotes (Sox2<sup>EGFP/IR</sup> and Sox2<sup>EGFP/LP</sup>) display a range of eye phenotypes from mild bilateral microphthalmia to severe anophthalmia. Histological analyses of postnatal Sox2<sup>EGFP/IR</sup> mice show that the majority of mutant retinas are ~30%–40% thinner than wild-type retinas. The thinning of the retina is associated with a reduction in cell number resulting in the decreased widths of the inner nuclear and plexiform layers (Fig. 3E, F). Additionally, a majority of mutant eyes display areas with disrupted cell layering, consisting of rosette structures (Fig. 3G, Fig. 6D–F, [below]). By E14.5, eyes in embryos of both Sox2 hypomorphic/null compound heterozygotes (Sox2<sup>EGFP/IR</sup> and Sox2<sup>EGFP/LP</sup>) are smaller than eyes of Sox2 heterozygous (Sox2<sup>+/EGFP</sup>) or wild-type littermates (Sox2<sup>++/+</sup>) (Fig. 3H, I; Supplementary Fig. 1) and, consistent with a dose-dependent function for SOX2, complete ablation of SOX2 specifically in retinal progenitors (aP0-CRE;Sox2<sup>EGFP/COND</sup>) correlates with a further reduction in eye size (Fig. 3J, see below).

SOX2 is required for the proliferative and differentiation capacity of retinal progenitors

To determine the requirement for SOX2 in retinal progenitor cells, we used a Cre-loxP strategy to conditionally inactivate Sox2 specifically in the mouse neural retina. In aP0-CREiresGFP mice, the expression of CRE and GFP is regulated by a retina-specific mouse Pax6 regulatory element such that CRE-recombinase activity is detected exclusively in the neural retina from E10.5 (Marquardt et al. 2001). Consistent with the general re-

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**Figure 2.** Generation of allelic series of mouse Sox2 locus. (A) Targeting vector introduced two loxP sites flanking the Sox2 promoter and mRNA coding regions. Homologous recombination of this vector at the Sox2 genomic locus in ES cells resulted in the generation of two alleles. Complete insertion of the vector generated the Sox2<sup>COND</sup> allele, while partial incorporation of the 3′ part of this vector generated the Sox2<sup>2LP</sup> allele, which contains only the 3′ insertion, but not the 5′ loxP site. CRE-driven recombination in Sox2<sup>COND/COND</sup>ACTB-Cre mice (in ACTB mice CRE recombinase is expressed in the female germline) resulted in deletion of the Sox2 promoter and mRNA coding region as well as the Neo cassette, generating a null allele [Sox2<sup>COND</sup>]. (B) The DNA recombination events were confirmed by Southern blot analysis of mouse tail DNA. (C) In targeting vector B, the 3′ UTR of Sox2 was replaced by internal ribosome entry site [IREs] and dsRed2 coding sequence followed by a Neo cassette flanked with two loxP sites. The incorporation of the insertion into Sox2 locus generated the Sox2<sup>2IR</sup> allele. The restriction enzymes used were AvrII [A], NheI [N], EcoRI [RI], Sall [S], Sple [Sp], EcoRV [V]. (D) Segregation of the alleles in the mouse lines was confirmed by Southern blotting with the P1 after EcoRI digest. (E) Quantification of pups and embryos of each genotype recovered from indicated breedings.

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Figure 3. Analysis of Sox2<sup>+/−</sup> and Sox2<sup>+/+</sup> hypomorphic alleles. (A,C) Immunoblot of protein extracts from wild-type and Sox2 mutant embryos, E14.5 brains and eyes, respectively, were developed with antibodies against SOX2. β-ACTIN antibody was used as a loading quantity control and an antibody against EGFP was used to control for the amount of putative SOX2 expressing cells in the sample. Note that the expression level of EGFP from the Sox2 locus is preserved throughout. (B,D) Quantification of the immunoblot results. SOX2 expression in Sox2<sup>+/−</sup> was taken as baseline (100%), and EGFP expression in Sox2<sup>+/−</sup> was designated as 50%. N = 2 for each genotype. (E–G) Compared with the Sox2 heterozygous mouse (Sox2<sup>+/EGFP</sup>; E), the compound null hypomorphic adult (Sox2<sup>+/EGFP</sup>; F) shows a significant reduction in the thickness of the retina. By E14.5, the null hypomorph exhibits distinct rosette structures (Sox2<sup>+/EGFP</sup>; G). (H–J) Comparison of eye morphology of E14.5 Sox2<sup>+/EGFP</sup>, Sox2<sup>+/+/EGFP</sup>, and Sox2<sup>+/−</sup>EGFP/IR, and Sox2<sup>+/−</sup>P0-CRE;Sox2<sup>+/−</sup>COND/+ mice. Bars: E,F, 50 µm; G, 100 µm; H–J, 200 µm.

A hallmark of SOX2-expressing neural progenitors is the maintenance of proliferative and differentiation capacity. It was shown previously that inhibition of SOX2 signaling by overexpression of a dominant interfering form of SOX2 in chick neural progenitors results in their premature exit from the cell cycle (Bylund et al. 2003; Graham et al. 2003). By E13.5, retinas of Sox2<sup>−/−</sup>;αP0-CRE mice were crossed to Sox2<sup>++/+EGFP</sup>;αP0-CRE and Sox2<sup>−/−</sup>;αP0-CRE, SOX2 is eliminated from these cells (Fig. 4A,F and B,G, respectively).

A hallmark of SOX2-expressing neural progenitors is the maintenance of proliferative and differentiation capacity. It was shown previously that inhibition of SOX2 signaling by overexpression of a dominant interfering form of SOX2 in chick neural progenitors results in their premature exit from the cell cycle (Bylund et al. 2003; Graham et al. 2003). By E13.5, retinas of Sox2<sup>−/−</sup>;αP0-CRE embryos (Fig. 4H) are significantly smaller in size compared with Sox2<sup>−/−</sup>;αP0-CRE retinas (Fig. 4C) as shown by PCNA staining. To further determine whether the hypocellular phenotype observed in Sox2<sup>−/−</sup>;αP0-CRE retinas is due to decreased proliferation, we assayed the expression of PCNA and incorporation of Bromodeoxy-Uridine (BrdU) into S phase nuclei. Although PNCA (Fig. 4H) remained expressed throughout the distal-central retina (Fig. 4D), there is a marked reduction of BrdU incorporation and exit from cell cycle in Sox2<sup>−/−</sup>;αP0-CRE embryos compared with cells of the central retina, thus indicating exit from cell cycle (Fig. 4I). In contrast, BrdU labeling in control embryos (Sox2<sup>−/−</sup>;αP0-CRE) is evenly distributed throughout the distal-central retina (Fig. 4D). Furthermore, the DNA content of the distal retina (GFP positive) cells of wild-type (Sox2<sup>+/−</sup>;αP0-CRE) and mutant Sox2<sup>−/−</sup>;αP0-CRE mice was compared using flowcytometric cell cycle analysis. These data show that ablation of SOX2, specifically in the cells expressing αP0-CRE, decreases the number of cycling cells (S/G2/M) from 31% to 8% (Supplementary Fig. 2). In addition to a decrease in BrdU incorporation and exit from cell cycle, Sox2 mutant distal RPCs, while maintaining expression of PCNA and Pax6, do not express molecular markers that define undifferentiated cycling RPCs such as Notch1 (Fig. 4E) and Hes-5 (Fig. 4K,P). Together these results demonstrate that SOX2 is required for appropriate neural retinal progenitor proliferation and maintenance of molecular markers that define progenitor identity.

To determine whether SOX2-null retinal cells that exit cell cycle differentiate into post-mitotic neurons, we assayed the expression of neuronal markers Math5, NeuroD, βTUBULINIII (Fig. 4L–N,Q–S), and ISLET1 (data not shown). In distal regions of Sox2<sup>−/−</sup>;αP0-CRE retinas, Math5, NeuroD, and βTUBULINIII were absent (Fig. 4Q–S). Thus, complete genetic ablation...
of Sox2 results in the loss of general retinal progenitor cell characteristics: the ability to proliferate and appropriately differentiate. Consistent with previous reports [Kammandel et al. 1999], we detected a variegated expression of the CREiresGFP transgene driven by the combination of the α enhancer and P0 promoter of
mouse Pax6 gene within the distal retina. The observed variability was increased in Sox2<sup>α0P-CRE</sup> mice ranging from mosaic CREiresEGFP expression restricted to the distal retina to widespread CREiresEGFP distribution throughout the circumference of the retina (Supplementary Fig. 3, cf. A and D). The coincident ablation of SOX2 with expression of CRE and the mutually exclusive expression of neuronal differentiation marker βTUBULINIII (red) and CREiresEGFP (green) in Sox2<sup>α0P-CRE</sup> embryonic retinas (Fig. 4S; Supplementary Fig. 3) indicates that SOX2 acts in a cell autonomous manner. A cell-autonomous role for SOX2 in retinal neural progenitor cells could explain the varying degrees of retinal cell loss in Sox2<sup>α0P-CRE</sup> (Fig. 4T) compared with Sox2<sup>+/α0P-CRE</sup> (Fig. 4O).

These results show that SOX2 plays a role in maintaining retinal neural progenitors in an undifferentiated proliferative state and, taken together with results from studies of SOX2 in the developing neural tube, imply a general requirement for SOX2 in neural progenitor cell populations.

**Eye morphogenesis in Sox2 hypomorphic mice**

The progressive increase in the severity of eye phenotypes displayed by mice carrying Sox2 mutant alleles (Fig. 3H–J) strongly suggests that the level of SOX2 is critical for normal retinal development. It is becoming recognized that the absolute level of a specific transcription factor is an important component of the mechanism of lineage specific regulation (Kulessa et al. 1995; DeKoter and Singh 2000; Motohashi et al. 2000, 2004; Emambokus et al. 2003). Moreover, it has been proposed that neural progenitor cell heterogeneity is triggered in part through a gene-dosage effect (Marquardt and Gruss 2002; Sinor and Lillien 2004).

To directly determine the functional significance of decreasing levels of SOX2 expression, we analyzed the phenotypic and molecular consequences in retinal neural progenitors of Sox2 hypomorphic/null mouse mutants [Sox2<sup>EGFP/IR</sup> and Sox2<sup>EGFP/LP</sup>]. Gross morphological analyses of Sox2<sup>EGFP/IR</sup> brains show hypoplasia of optic nerves and chias mata (Fig. 5A,E). RGCs are the only retinal projection neurons to extend axons outside the eye; their axons grow laterally toward the presumptive optic nerve head and pass outward through the optic stalk to travel along the optic nerve to the chiasm. Thus, the observed failure to form optic nerves in Sox2<sup>EGFP/IR</sup> mice strongly suggests a defect in RGCs due to early disruption of SOX2.

To determine whether the Sox2<sup>IR</sup> mutation affects differentiation of retinal neurons, we used immunostaining to evaluate the differentiation of distinct cell types in postnatal Sox2<sup>EGFP/IR</sup> retinas. RGCs, as marked by NEUROFILAMENT [Fig. 5B,F, arrowhead] and horizontal cells [NEUROFILAMENT, Fig. 5I,M, arrowheads], bipolar cells [PKC, Fig. 5J,N], rod photoreceptors [RHODOPSIN, Fig. 5K,O, arrowhead] and horizontal cells [NEUROFILAMENT, Fig. 5B,F, arrowhead] are expressed, and Müller glia are formed in Sox2<sup>EGFP/IR</sup> retinas (GFAP, Fig. 5L,P; CRABLP, Supplementary Fig. 4D,F). Interestingly, whereas GFAP expression in the wild-type and heterozygous (Sox2<sup>+/GFP</sup>) mouse retina is restricted to the distal region, the Sox2<sup>EGFP/IR</sup> retina expresses GFAP throughout the distal–central axis. Colabeling with CRABLP identified these cells as Müller glia (Supplementary Fig. 4), which in the central region of the wild-type mouse retina do not normally express GFAP. Therefore, the observed increase in GFAP expression in the Sox2<sup>EGFP/IR</sup> retina appears to be an up-regulation of GFAP in these cells and not an increase in their cell number. In addition to an apparent absence of RGCs, the majority of Sox2<sup>EGFP/IR</sup> eyes show disrupted cell layering, which is demonstrated by the presence of rosette structures (Figs. 3, 6 [cf. A–C and D–F]). The rosette structures are comprised of differentiated retinal neuronal subtypes. For example, they contain rod cells (marked by RHODOPSIN, Fig. 6C,F) surrounded by horizontal cells (marked by NEUROFILAMENT, Fig. 6B,E) and amacrine cells (marked by CALRETININ, Fig. 6A,D).

**Reduction in SOX2 levels restricts retinal neural progenitor competence**

To determine whether RGCs are generated during embryogenesis in Sox2<sup>EGFP/IR</sup> mice, we assessed Brn3b and βTUBULINIII expression, which at E15 identify differentiating RGCs. In both the wild-type and Sox2<sup>EGFP/IR</sup> mutant retina, Brn3b and βTUBULINIII-positive cells could be identified in the inner side of the developing neural retina, consistent with the formation of RGCs (Fig. 6G,L,H,M). Although RGCs were present in the Sox2<sup>EGFP/IR</sup> mutant retina, the cells were inappropriately localized. For example, βTUBULINIII-positive cells were not restricted to the inner side, but could also be detected in the outer RBL layer, which normally at this stage consists of undifferentiated progenitor cells (Fig. 6M, arrows). These regions contained fewer progenitors, as marked by PCNA, and showed disruption of retinal laminar structure (Fig. 6I,N, arrow). Moreover, the RGC axons of Sox2<sup>EGFP/IR</sup> retinas, as marked by βTUBULINIII [Fig. 6, cf. H and M] and anterograde labeling of retinal ganglion cells with Dextran-Cy3 (Fig. 6, cf. J and O, dotted outline), are scattered to subretinal spaces and fail to enter the optic nerve. This failure of SOX2 hypomorphic RGCs to reach their target is consistent with an increase in apoptotic cells as indicated by Caspase 3 staining (Fig. 6K,P).

Neural progenitor cells are characterized and defined by their pseudopodial morphology, proliferative capacity, and gene-expression profiles. The disruption in retinal structure and aberrant ganglion cell differentiation in Sox2<sup>EGFP/IR</sup> retinas indicates a change in progenitor properties due to a decrease in the level of SOX2. To address this question and begin to determine the molecular basis for these defects, we examined the expression of markers that define progenitor cells and the onset...
of their differentiation in Sox2EGFP/IR mice, focusing on regulators of RGC differentiation. A number of transcription factors have been shown to serve as intrinsic regulators of RGC differentiation, among them the homeodomain factor PAX6 and basic helix–loop–helix (bHLH) factors MATH5, NEUROD, HES-1, and HES-5, GLI1 (Marquardt and Gruss 2002; Hatakeyama and Kageyama 2004; Mu et al. 2005). PAX6 is required for the activation of MATH5, an essential factor for RGC lineage specification. NEUROD1, another member of bHLH proneural genes, is also implicated in promoting retinal neuronal differentiation. In contrast, the NOTCH pathway negatively regulates RGC differentiation by activating the Hairy-enhancer of Split [HES] family of bHLH genes, Hes-1 and Hes-5, thereby repressing the proneural bHLH factor Math5. Thus, the PAX6 and NOTCH pathways counter each other in the regulation of downstream genes required for the generation of the RGC lineage. To assess the pathways disrupted in Sox2 mutants, we therefore compared Math5, NeuroD1, Glil, PAX6, Notch1, and Hes-5 expression between the wild-type and Sox2 hypomorphic/null mutant retina at E14. In both wild-type and Sox2EGFP/IR retinas, Math5, NeuroD, Glil, and PAX6, are expressed at high levels in the retinal outer layer [Fig. 7, cf. A–C,G,D–F] consistent with progenitor cell differentiation and generation of RGCs. In contrast, expression of Notch1 and Hes-5 [Fig. 7, cf. H,I and K,L], which function to repress RGC dif-

Figure 5. Decrease in SOX2 levels leads to hypoplasia of the optic nerve and loss of retinal ganglion cells. Ventral side of 3-mo-old Sox2+/EGFP [A] and Sox2EGFP/IR [E] brains. [E] Note the absence of developed optic nerve [arrow] in Sox2EGFP/IR. Immunostained sections through the retina of adult Sox2+/EGFP [B–D,I–L] and Sox2EGFP/IR [F–H,M–P] mice. Compared with the Sox2+/EGFP, Sox2EGFP/IR mice show ~95% reduction in the number of cells immunostained for NEUROFILAMENT [B,F, arrows], Bm3b [C,G, arrows], and ISLET1 [D,H, arrows], and a ~60%–70% reduction in cells stained with CALRETININ [I,M, arrowhead] in the ganglion cell layer [GCL]. In contrast, interneurons demonstrate a relatively normal distribution, as shown by staining for NEUROFILAMENT [B,F, top arrowheads], ISLET1 [D,H], CALRETININ [I,M], and PKC [J,N]. [I,M] Note the significant reduction in size of inner plexiform synaptic layer [IPL] as visualized by CALRETININ staining [arrowheads]. [K,O] No cell loss was apparent in outer nuclear layer (ONL) formed by photoreceptor cell bodies, as confirmed by RHODOPSIN staining of the outer segments [arrowheads]. [L,P] We noted an up-regulation of GFAP expression in central retina of the Sox2EGFP/IR mice as compared with the Sox2+/EGFP mice [insets, gross morphology]. Bars: A,E, 2 µm; B–D,F–P, 100 µm; L,P [insets], 200 µm.
differentiation and maintain progenitor identity, are significantly down-regulated in retinal progenitors of Sox2 hypomorphs. Wild-type levels of Notch1 and Hes-5 expression are maintained in the neural tube where SOX2 is coexpressed with SOX1 and SOX3, providing an internal control for in situ staining (Supplementary Fig. 5A–D). Furthermore, Western blot analyses of SOX2 and NOTCH1 expression in retinal cells isolated from wild-type, heterozygous, and hypomorphic/null Sox2 retinas indicate that the decrease in levels of SOX2 expression directly correlates with the decrease in levels of NOTCH1 expression (Supplementary Fig. 5E).

In Sox2EGFP/IR mutant retinas, down-regulation of SOX2 results in a decrease in the level of expression of NOTCH1 and its direct downstream target Hes-5, while expression of PAX6 and Math5 are up-regulated and Gli1 and NeuroD remain expressed. These data indicate that SOX2 may maintain retinal progenitor identity, at least in part, through the regulation of NOTCH1 signaling.

SOX2 transcriptionally regulates Notch1 expression

To establish whether Notch1 is directly regulated by SOX2, and to elucidate the molecular mechanisms un-
derlying SOX2 function, we conducted a chromatin immunoprecipitation (ChIP) cloning screen—a method that allows for the identification of direct downstream targets of a transcription factor [Weinmann et al. 2002]. To this end, a library of ChIP DNA fragments was generated from embryonic stem (ES) cells using a SOX2 polyclonal antibody [Fig. 8A]. Fifty clones were sequenced, and BLASTed against the whole mouse genome to identify the loci of the ChIP DNA fragments. Among the loci identified was \textit{Fgf4}, a well-characterized direct target of SOX2 [Dailey et al. 1994]. More critically we also identified a DNA fragment localized to intron 13 (IVS13) of the \textit{Notch1} locus (Fig. 8C).

To further validate the in situ binding of SOX2 to the IVS13 region of \textit{Notch1}, and to demonstrate that the same SOX2-dependent enhancer of \textit{Notch1} could be ChIPed from CNS tissue, a ChIP PCR analysis [Fig. 8B] was conducted using mouse ES cells, E14.5 CNS tissue, and mouse embryonic fibroblasts (MEF, which lack SOX2), as a negative control. The results show that IVS13 of \textit{Notch1} is enriched upon ChIP in both ES cells and E14.5 CNS, but not in MEFs, or when using a non-specific IgG antibody [Fig. 8C].

The 2.0-kb \textit{Notch1} IVS13 DNA fragment identified by ChIP was assessed for putative SOX2-binding sites. Bioinformatic analysis identified three SOX consensus sites within the 2.0-kb DNA fragment [Fig. 8D]. To confirm SOX2 binding to the putative \textit{cis} elements, DNase I footprinting analysis was conducted using a 150-base-pair (bp) probe that contained all three sites. Using HIS-tagged purified SOX2 protein, three strong DNase I interference patterns were observed, which directly correlate with the locations of the three SOX consensus sequences [Fig. 8E].

Furthermore, to determine whether the 150-bp fragment had the ability to enhance transcription, the \textit{Notch1} IVS13 fragment containing the SOX2 consensus sites was cloned into a minimal promoter luciferase reporter gene construct [Tugores et al. 1994] that was transiently transfected into the early ectodermal cell line P19 [Fig. 8F]. The results show that the 150-bp \textit{Notch1} IVS13 DNA fragment increases transcription compared with the minimal promoter alone. Moreover, site-directed mutagenesis of the +26988 and +27001 SOX2 \textit{cis} elements ablates any enhanced transcription, indicating that the SOX2 consensus binding sites are critical for the transcriptional enhancement of the reporter gene construct. Together, these data strongly imply that \textit{Notch1} is transcriptionally regulated by SOX2.

**Discussion**

In this study we provide genetic evidence demonstrating that alterations in the levels of SOX2 regulate the choice between maintenance of retinal progenitor cell identity and differentiation. Wild-type levels of SOX2 maintain progenitor developmental potency, the ability to proliferate and differentiate, while a decrease in SOX2 levels (<40% of wild-type levels) results in the restriction of progenitor competence including the disruption of neuroepithelial morphology, proliferative capacity, and aberrant differentiation. In part, these characteristics of de-
SOX2 and retinal progenitor fate

Figure 8. SOX2 binds IVS13 cis elements in the Notch1 gene. (A) Whole-cell extracts from ES cells and Control STO cells (which lack SOX2) were incubated with SOX2 antibody. SOX2 was efficiently immunoprecipitated (IPed) from only the ES cell extracts. [B] Schematic representation of standard ChIP PCR assay. PCR primers flanking the SOX2 consensus cis elements were used in a standard ChIP PCR assay with DNA template IPed from ES cells, E14.5 CNS, or MEFs as a control. Sonicated DNA was volumetrically divided in half to ensure equivalent input templates, and ChIP was conducted with either a SOX2 or nonspecific IgG as a control antibody. [C] The Notch1 IVS13 locus was IPed as a direct function of SOX2 binding in both ES cells and E14.5 CNS, but not from MEFs [which lack SOX2]. Known SOX2 cis elements positioned in well-characterized enhancers of Fgf4 and Nestin were used as positive controls to demonstrate efficient ChIP using the SOX2 antibody. [D] Probe sequence used for DNase I footprinting analysis. Numbers indicate the nucleotide positions of the SOX2 consensus sites within Notch1. The underlined sequence represents the area of the footprint pattern. Red boxes indicate bases in the SOX2 consensus sites that were mutated (+26988: TGT to CC, and +27001: TGT to GCA). [E] DNase I footprinting demonstrates the binding of proteins that produce interference patterns that localize specifically to SOX2 consensus sites within the Notch1 IVS13. Numbered arrows on the left indicate the location of the SOX2 consensus sites in the probe. The schematic diagram on the right represents the approximate locations of SOX2 binding to the probe. [F, left schematic] Luciferase reporter gene constructs were generated that contained the intact 150-bp IVS13 DNA fragment or the same DNA fragment in which either the +26988 or +27001 SOX2 consensus sites were mutated. [Right graph] Transient transfections in P19 cells indicate that the 150-bp IVS13 of Notch1 has the ability to enhance transcription of the reporter gene by approximately threefold, and that the +26988 and +27001 SOX2 consensus sites are required for transcriptional enhancement.

Effective retinal development in a Sox2-deficient genetic background are a consequence of altered NOTCH1 signaling. Our study shows a direct relationship between SOX2 and the Notch1 gene in that SOX2 interacts directly with a Notch1 enhancer, and that, in turn, the levels of SOX2 directly correlate with the levels of NOTCH1.

A role for SOX2 in vertebrate eye development was first suggested based on its expression pattern; SOX2 is expressed throughout the cells of optic vesicle of the anterior neural plate, transiently expressed in the newly induced lens, and is then restricted to proliferating progenitor cells of the neural retina [Kamachi et al. 1998; Van Raay et al. 2005]. Consistent with its early expression in the anterior neural plate, inhibition of SOX2 signaling by injection of antisense morpholinos or dominant interfering forms of Sox2 into Xenopus embryos results, in addition to other morphological effects, in reduced or missing eyes [Mizuseki et al. 1998; Van Raay et al. 2005]. Furthermore, misexpression of SOX2 in chick embryos has been shown to induce ectopic lens formation as marked by β-CRYSTALLIN [Kamachi et al. 2001]. In order to bypass the early requirement of SOX2 in the anterior neural plate and to specifically address its role in neural retinal progenitor cells, we used the Cre-loxP strategy to conditionally inactivate Sox2 in the mouse. Our results demonstrate that SOX2 is required for the maintenance of neural retinal progenitor identity, such that after Cre-mediated ablation of SOX2 activity in the developing neuroretina, RPCs lose their ability to both divide and differentiate. Consequently, neurogenesis, including the up-regulation of bHLH factors [MATH5, NEUROD1] and the appearance of βTUBULINIII differentiated neurons, was not observed in the Sox2A/CND/ACND; aP0-CRE mutant retinas. The cellular and molecular consequences of genetic ablation of Sox2 in progenitor cells of the neural retina in the mouse and the morpholino-knockdown of SOX2 in Xenopus further highlight an evolutionarily conserved function and requirement of SOX2 in retinal progenitor cell biology [Van Raay et al. 2005].

The complete genetic ablation of Sox2 described here in retinal neural progenitor cells phenocopies inhibition of SOX2 signaling in embryonic and adult neural tube progenitors. First, expression of a dominant interfering form of SOX2 in chick neural progenitors results in their premature exit from cell cycle [Bylund et al. 2003; Graham et al. 2003]. Second, the conversion of rat oligodendrocyte precursors [OPCs] into multipotent neural stem cells [NSCL] is dependent on reinitiation of SOX2 expression, and inhibition of SOX2 signaling results in premature exit from cell cycle and neuronal differentiation of OPCs [Kondo and Raff 2004]. Finally, it has recently been demonstrated that SOX2 expression is maintained in neurogenic regions of the adult rodent nervous system and that regulatory mutations of mouse Sox2 cause neurodegeneration and impair adult neurogenesis [Ferri et al. 2004]. The conserved requirement for SOX2 in the CNS ventricular zone and retinal progenitors suggests that a common SOX2-regulated mechanism regulates the
maintenance of progenitor cell identity in distinct regions of the CNS.

Despite mounting evidence that SOX2 is important in the maintenance of progenitor cell identity (Bylund et al. 2003; Graham et al. 2003; Ferri et al. 2004), the question remains: Does SOX2 interact with other genetic pathways implicated in this process, and if so, how? The maintenance of CNS progenitor identity is critically dependent on NOTCH signaling. NOTCH1, along with its two direct downstream targets HES-1 and HES-5, is co-expressed in the ventricular zone of the developing central nervous system including the retina, and maintains RPCs in an uncommitted state [Hatakeyama and Kageyama 2004; Hatakeyama et al. 2004]. Moreover, the transition from uncommitted to lineage-restricted RPCs is coupled with both the down-regulation of NOTCH1 activity, presumably in response to extrinsic signals, and the activation of retinogenic bHLH factors such as MATH5, NEUROD1, and NGNs. However, what triggers the transition from an “uncommitted RPC” to a “lineage-restricted RPC” still remains unclear; for example, what intrinsic differences occur in RPCs when challenged with the same extrinsic signals?

Restriction of neural progenitor cell capacity is directly coupled with the down-regulation of SOX2 expression. Heterogeneity in SOX2 levels among progenitors could therefore allow cells to respond differently to the same extrinsic signals. By generating an allelic series of Sox2 to directly analyze the effects of graded reduction of SOX2, we provide genetic evidence that one of the intrinsic differences in progenitor cells is the level of SOX2, which may act by modulating NOTCH1 expression in neural progenitors. We present three independent sets of experiments demonstrating that Notch1 transcription is regulated by SOX2, an unbiased ChIP screen identified Notch1 as a direct target of SOX2 in vivo, DNase footprinting and Luciferase reporter assays showed that SOX2 can bind to Notch1 regulatory regions and regulate the levels of NOTCH1 expression in vitro, and finally, both NOTCH1 and its target Hes-5 are dramatically down-regulated in the retinas of Sox2 hypomorphic mice. Furthermore, ablation of NOTCH1 gives a retinal phenotype similar to that which we describe for Sox2-null mice [i.e., loss of RPCs] (Austin et al. 1995; Waid and McLoon 1995; Ahmad et al. 1997; Henrique et al. 1997; Jadhav et al. 2006], and the retinal phenotypes of both Hes-1 and Hes-5 mutant mice are strikingly similar to the phenotype observed in Sox2-hypomorphic mice, in that RPCs aberrantly differentiate and abnormal “rosette-like” structures are present [Ishibashi et al. 1994; Tomita et al. 1996]. The lack of balance between lateral inhibitory signaling by NOTCH1 and proneural gene products results in differentiation of RPCs—a mechanism that may extend to other CNS progenitors. The data presented in this study provide a molecular explanation for the inverse correlation between the levels of proneural bHLH and SOXB1 factor expression in chick spinal-cord progenitor cells in that the capacity of proneural bHLH proteins to direct neuronal differentiation crucially depends on the suppression of SOX1, SOX2, and SOX3 (Bylund et al. 2003). In light of the conserved expression/function of NOTCH1 signaling in other stem cell/progenitor populations, and the ability of other SOX proteins to bind the loose SOX consensus sequence, other SOX family members may act in a similar manner to universally conserve the “stemness” identity. Consistent with this, SOX10 has been implicated to act in a dosage-dependent manner in the peripheral nervous system to maintain neural crest stem cell identity by indirectly regulating the expression of bHLH factors [Kim et al. 2003].

Our data show that cell fate determination of retinal progenitor cells is dependent on the levels of SOX2, however, the molecular mechanism by which SOX2 dosage regulates cell fate remains to be elucidated. To date, two major mechanisms of transcriptional regulation by SOX factors have been suggested, protein–protein interaction and modification of chromatin structure [Pevny and Lovell-Badge 1997; Kamachi et al. 1999]. The expression of FGFR4 and NESTIN, both previously identified as direct transcriptional targets of SOX2, is dependent on heterodimerization of SOX2 with protein partners, OCT3/4 and Brn2, respectively [Tanaka et al. 2004]. In the context of an interacting protein network, it is possible that subtle differences in protein levels are sufficient to shift the equilibrium toward another transcription complex. Furthermore, binding of SOX factors, including SOX2, change the local conformation of DNA, i.e., introducing or stabilizing bends that may determine the accessibility of DNA to other transcription complexes [Scalfieri and Bianchi 2001]. It is tempting to speculate that transcriptional networks, such as those including SOX factors, are highly sensitive to a dosage effect that can modify their capacity to form particular protein–protein interactions and/or affect changes in higher order chromatin structures. Precedent for this comes from studies of mechanisms regulating hematopoietic stem cell differentiation. For example, a gradient of the E2A transcription factor has been shown to determine the choice between stem, B- or T-cell fate. Moreover, based on NOTCH1 signaling studies, a reduction in E2A function in the common lymphoid precursor is permissive for T-cell lineage but not for the B-cell lineage that requires intact E2A activity [Zhuang et al. 2004].

The critical importance of precise regulation of transcription-factor dosage is further reinforced by the genetics of human syndromes. Humans with mutations in SOX2 display a spectrum of severe eye malformations ranging from unilateral microphthalmia to bilateral anophthalmia including, in some cases, aplasia of the optic nerve, chiasm, and optic tract. The severity of the disease state most likely reflects genetic variations within the human population, or alternatively, reflects the strength of the SOX2 mutation (e.g., null vs. hypomor- phic). Detailed analyses of the cellular defects and molecular pathways associated with human SOX2-related retinal defects have been difficult to assess in patients, since the onset of the defect is developmental. Therefore, a viable approach to circumvent this problem is to generate a mouse model in which the earliest phenotypic
sequelae of aberrant SOX2 expression can be monitored and studied in the embryo. Our analyses of an allelic series of Sox2 mutations in the mouse have confirmed a critical relationship between gene dosage of Sox2 and the severity of eye malformations. Microphthalmic and optic nerve hypoplasia was observed in Sox2<sup>COND/COND</sup>;αP0-<SUP>CRE</SUP> animals, thus implicating the disruption of SOX2 function specifically in the neural retina as a contributing factor to the phenotype observed in human SOX2-Anophthalmia. Compound hypomorphic null mice (Sox2<sup>EGFP/LP</sup>, Sox2<sup>EGFP/IR</sup>) display a series of eye malformations, and in most severe cases, both eyes fail to develop (bilateral anophthalmia). Moreover, we have demonstrated that these differences in the severity of eye phenotypes correlate with the defects in the neural retina, and they are directly related to the levels of SOX2 expression. Although the phenotype observed in Sox2<sup>-</sup>hypo- morphic mice [compound heterozygotes] closely mimics that displayed in humans with Sox2 mutations, the defect in humans is primarily defined as a heterozygous disorder where one allele possesses a missense or null mutation, and the other allele is “wild type.” Although we cannot rule out that species differences may account for a phenotypic difference between mouse and human, an intriguing point of speculation based on the genetic consequences of the Sox2<sup>-</sup>-hypomorphic mouse is that humans with Sox2-related retinal defects may indeed be compound heterozygotes; that is, they may possess a null allele and a “low-expressing” allele that brings the total functional level of SOX2 to phenotypic levels (<40% of normal). The literature is replete with examples of human disorders in which a low-expressing allele, combined with a null-allele, precipitates the disease condition. The molecular defects associated with a “low-expressing allele” range from enhancer defects to aberrant splicing and mRNA stability. Our study may shed light on potential regulatory mutations in the “unexplored” allele from cohorts already identified to possess a null-SOX2 allele. An alternative hypothesis is that humans with SOX2-related eye disorders have epistatic mutations or complex genetic interactions that result in retinal and/or lens defects. The Sox2<sup>EGFP</sup> and Sox2<sup>-</sup>-hypomorphic mice provide critical genetic tools to identify interacting pathways necessary for normal eye development and causative to pathologic conditions.

**Materials and methods**

**Gene targeting and mouse breeding**

The Sox2<sup>COND</sup> targeting construct was generated by inserting a loxP site in the SpeI site 5′ of the first Sox2 exon and a PGK–Neo cassette flanked by a loxP site in the Nhel site immediately 3′ to the Sox2 coding region (Avilion et al. 2003). The Sox2<sup>-</sup>-targeting construct was created by partial incorporation of the Sox2<sup>COND</sup> targeting vector into Sox2 locus. This introduced the PGK–Neo cassette with a single loxP site into the first Nhel site 3′ of Sox2 coding sequence (Fig. 2A). In the Sox2<sup>-</sup>-construct, the 3′UTR of Sox2 mRNA (EcoRV–Nhel fragment) was replaced with IRES<sup>dlsRed2</sup> (Clontech) and loxP-flanked PGK–Neo cassette (Fig. 2C). Targeted constructs were electroporated into CCE ES cells [gift from E. Robertson, Oxford University, Oxford, UK] and recombinants were identified using standard Southern blot analysis with α<sup>32</sup>PdCTP DNA probes [Fig. 2B,D]. Chimeric founder mice that transmitted the targeted alleles were produced by blastocyst injection. Sox2<sup>COND/COND</sup> mice were generated by mating heterozygous Sox2<sup>COND</sup> mice to ACTB-Cre FVB/N-Tg(ACTB-cre)2Mrt/J, in which CRE recombinase is expressed in the female germline [Jackson Laboratories] (Lewan- doski et al. 1997) and αP0-Cre mouse line [gift from Dr. P. Gruss, Max-Planck-Institute of Biophysical Chemistry, Germany] (Marquardt et al. 2001). PCR genotyping of Sox2 alleles was performed using the following sets of primers: for the wild-type allele [Sox2<sup>+</sup>] 5′-GCTCTGGTTATGGGAATCAGGC-3′ and 5′-CTGCTCACGGAAAGGAGG-3′ primers; 35 cycles at 94°C for 15 sec, at 55°C for 30 sec, at 72°C for 30 sec; generated 382-bp product, for targeted alleles [Sox2<sup>EGFP</sup>, Sox2<sup>Lp</sup> and Sox2<sup>-</sup>] 5′-CACAGGCTCTTCATGCACTACAG-3′ and 5′-CAACGATTTTCAGGTTC-3′ primers; 35 cycles at 94°C for 15 sec, at 56°C for 30 sec, at 72°C for 40 sec; generated 297-bp product. The Sox2<sup>COND</sup> allele was detected using 5′-CTTTTCCTGTTGTGATGCTTTCG-3′ and 5′-ATCTGTTGTGCCTGGAACAGTATCC-3′ primers; 45 cycles at 94°C for 15 sec, at 55°C for 30 sec, at 72°C for 40 sec; by 589-bp product. Sox2<sup>EGFP</sup> allele [Ellis et al. 2004] was genotyped using 5′-CGCTTTCTCTGTTCTTAC-3′ and 5′-GGCTTTCTCTTTTTCTCTCTTAG-3′ primers; 35 cycles at 94°C for 15 sec, at 60°C for 30 sec, at 72°C for 45 sec, by 750-bp product.

**Northern blot analysis**

Total RNA was extracted from E14 mouse brains using Trizol reagent [Invitrogen] and analyzed by Northern blotting using α<sup>32</sup>PdCTP DNA probes for Sox2 [PS2, Fig. 2A] and β-Actin [939-bp fragment generated using 5′-GTGACGAGGCCCAGAG primers; 35 cycles at 94°C for 15 sec, at 55°C for 30 sec, at 72°C for 40 sec; by 589-bp product. Sox2<sup>EGFP</sup> allele (Ellis et al. 2004) was genotyped using 5′-CGCTTTCTCTGTTCTTAC-3′ and 5′-GGCTTTCTCTTTTTCTCTCTTAG-3′ primers]. The blots were developed using a PhosphorImaging Typhoon scanner [Amersham Biosciences] and quantified using ImageQuant [Amersham Biosciences].

**Immunoblotting**

Protein extracts were prepared from E14 embryonic brains or eyes in RIPA buffer [Magnees et al. 1998] containing protease inhibitor cocktail [Sigma]. Immunoblotting was conducted according to standard protocol using primary antibodies against SOX2 [Chemicon; 1:1000], EGFP [Molecular Probes; 1:5000], NOTCH1 [Santa Cruz Biotechnology; 1:1000] and βACTIN [Sigma; 1:5000]. IgG-horseradish conjugate [Bio-Rad] was applied as the secondary antibody [1:10,000] and Enhanced Chemiluminescence Reagent [Amersham] was used for detection. Bands were quantified by scanning the film and the image densities analysis using NIH Image software [http://rsb.info.nih.gov/nih-image/Default.html].

**Tissue preparation, immunohistochemistry, and in situ hybridization**

Mouse embryos were fixed with 4% paraformaldehyde [PFA] in phosphate-buffered saline [PBS]. Adult mice were perfused with 4% PFA and the eyes and brains were post-fixed for 30 min at room temperature. Tissue was immersed sequentially in 10%, 20%, and 30% sucrose/PBS and then embedded and frozen in OCT medium [Tissue-Tek]. For BrdU labeling, pregnant mice were injected intraperitoneally with BrdU [100 mg per kilogram of body mass] and euthanized 2 h thereafter. Horizontal 12–14 µm cryostat sections were blocked in 1% goat serum in PBS.
0.1% Triton-X 100 and incubated with primary antibodies at 4°C overnight and with secondary antibodies for 20 min at room temperature. Working dilutions and sources of the following antibodies were used in this study: SOX2 (Chemicon, 1:2000), SOX1 (Pevny et al. 1998) (1:500), PCNA (Biosource International, 1:500), BrdU (Becton Dickinson, 1:100), GABRB3 (Covance, 1:1000), NEUROFLAMET (Hybridoma Bank, 1:5000), CALRBP (1:4000), GFAP (DAKO, 1:500), PAX6 (Hybridoma Bank, 1:50), PAX6 (Covance, 1:200), ISET1 (Hybridoma Bank, 1:50), CALRETNIN (Chemicon, 1:1000), Protein Kinase C α (Sigma, 1:1000), RHODOPSIN (Leinco Technologies, 1:50), CRE-recombinase (Novagen, 1:2000), activated CASPASE3 (Cell Signaling Technology, 1:50), Cy3-anti-mouse (Sigma, 1:200), Cy3-anti-rabbit (Sigma, 1:200). In situ hybridization was performed on 20-µm horizontal cryostat sections using DIG-labeled antisense RNA probes and followed by enzymatic detection according to manufacture protocols (Roche). The following probes were used: Math5 (Yang et al. 2003), NeuroD1 (Lee et al. 1995), Notch1 (Lardelli and Lendahl 1993), and Hes-5 (Chen and Walsh 2002).

**ChIP library and PCR analysis**

A SOX2 ES ChIP library was generated essentially as described in Weinmann and Farnham (2002). Anti-SOX2 antibody (Chemicon; 2.5 µg/ml) was used for IP with Protein A/G Agarose Plus (Santa Cruz Biotechnology) beads. ChIP material (1 µg) was used in a Zero-blunt TOPO cloning reaction (Invitrogen) with T4 DNA ligase and 100 ng of the plasmid pCAGGS vector. ChIP library and PCR analysis was carried out in duplicate, and the transfection efficiency of Notch1 IVS13 reporter genes was normalized to CMV-RENilla luciferase activity.

**Footprinting protection assay**

Full-length HIS-tagged SOX2 (pcDNA3.1- SOX2-V5HIS) was transiently expressed in 293 cells, purified with Ni-NTA beads according to manufacturer’s protocol (Qiagen), and dialyzed against Dignam C with 50 mM NaCl. Purification was assessed by Coomassie-stained SDS-PAGE gels and immunoblotting. A footprinting assay was conducted essentially as described (Tu-gores et al. 1994) using 30 µg of purified SOX2-V5-HIS, dC: dG, and a probe generated from the 150-bp amplicon recovered by PCR using the Notch1 primers described above.

**Luciferase assay**

The 150-bp DNA fragment [IVS13] used in the DNase I footprinting assay was cloned into the 3’ polylinker of p21LUC, which contains a minimal promoter driving expression of Pho- tinus pyralis luciferase. Site-directed mutagenesis was carried out on the SOX2 consensus sites, +26988 and +27001, located in the 150-bp Notch1 IVS13 DNA fragment. The mutations were induced with the PCR primers +26988FOR [5’-CAGAGGTCAGCACTGCTCAAGGTTGGACT-3’ and the complementary primer] or +27001FOR [5’-CATTGTGCAAGGTTGGCACATGCACTGCTCAAGGTTGGACT-3’ and the complementary primer] using a modified QuickChange method (Stratagene). The +26988 site was converted from ATTTTC to ATTCCTG and the +27001 site was converted from ATTTTC to ATTCCTG. Notch1 IVS13 luciferase reporter gene constructs (1 µg/40 wells), and CMV-RENilla luciferase (0.2 µg/well) as a normalization control, were transiently transfected into the P19 cell line using Lipofectamine Plus reagent (Invitrogen) in OptiMEM medium (Gibco). After 12 h of exposure to transfection medium, the medium was replaced with DMEM-H (Gibco) supplemented with standard concentrations of glutamine, penicillin, and streptomycin, and incubated for an additional 36 h. Forty-eight hours after transfection, the expression of both P. pyralis and Renilla Luciferase were assessed using the Dual Luciferase Reporter Assay System (Promega). All experiments were carried out in duplicate, and the transfection efficiency of Notch1 IVS13 reporter genes was normalized to CMV-RENilla luciferase activity.

**Fluorescence-activated cell sorting (FACS)**

Retinas from E13.5 wild-type (Sox2+/−;aPO-CRE) and mutant (Sox2+/−;aPO-CRE;aPO-CRE) dissociated as described in Polleux and Ghosh (2002). Live cells (106) were stained for 30 min at 37°C with Vibriant Violet dye (Invitrogen) and then subjected to FAC analysis using a CyAn flowcytometer (DAKO Cytomation, Glostrup). DNA content was determined specifically in aPO-CREiresGFP cells by gating on EGFP-positive cells and assessing Vibriant Violet fluorescence. DNA Content histograms were modeled for cell cycle parameters in 30,000 cells using MODFIT software (Verity).

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