sox4 And sox11 Function during Xenopus laevis Eye Development

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Abstract

SoxC genes are involved in many developmental processes such as cardiac, lymphoid, and bone development. The SoxC gene family is represented by Sox4, Sox11, and Sox12. Loss of either Sox4 or Sox11 function is lethal during mouse embryogenesis. Here, we demonstrate that sox4 and sox11 are strongly expressed in the developing eye, heart as well as brain in Xenopus laevis. Morpholino oligonucleotide mediated knock-down approaches in anterior neural tissue revealed that interference with either Sox4 or Sox11 function affects eye development. A detailed analysis demonstrated strong effects on eye size and retinal lamination. Neural induction was unaffected upon Sox4 or Sox11 MO injection and early eye field differentiation and cell proliferation were only mildly affected. Depletion of both genes, however, led independently to a significant increase in cell apoptosis in the eye. In summary, Sox4 and Sox11 are required for Xenopus visual system development.

Introduction

In Xenopus, eye development starts with the induction of the eye field in the anterior neural plate at the end of gastrulation. The eye anlage is characterized by the expression of eye specific marker genes such as rax, pax6, and six3. Loss of any of these genes leads to strong defects during early eye development [1]. In contrast, the overexpression of these factors can result in the formation of ectopic eyes [1]. During neurulation, the single eye field splits into two eye anlagen located on either side of the embryo, a process supported by the underlying prechordal mesoderm [2]. At the end of neurulation, two optic vesicles evaginate from the neural tube at the level of the prospective diencephalon towards the overlying ectoderm. Through the contact of the eye vesicle with the epidermis, the lens placode is thickened in the epidermis. Later, the distal half of the eye vesicles invaginates into the embryo thereby forming a bilayered optic cup.

The optic cup can further be subdivided into the thinner outer retinal pigmented epithelium (RPE) and the thicker neural retina in which six major neuronal cell types and glial cells develop. In the mature retina, different well structured layers can be distinguished: the outer nuclear layer (ONL) containing the cell bodies of cone and rod photoreceptors, the inner nuclear cell layer (INL) including the cell bodies of bipolar, horizontal, and amacrine interneurons, and the ganglion cell layer (GCL) containing the cell bodies of ganglion cells. Muller glial cells span all retinal layers.

The founding member of the Sox (sry-related box) gene superfamily of transcription factors was the male sex determination gene Sry (sex determining region Y). Sox proteins contain a single high mobility group (HMG) domain, which is involved in DNA binding. This domain of all Sox gene family members has an identity of more than 50% to the HMG box of the Sry gene. Currently, 20 SOX genes in human and mouse are known which are classified into ten groups, Sox4 to Sox10, based on sequence similarity and similar DNA binding properties [3]. Through protein-protein interactions, Sox proteins are also able to recruit additional proteins to DNA. They can act either as transcriptional activators or repressors depending on the cellular context and the interaction partners [4]. During development, Sox proteins are essential for many processes such as the regulation of pluripotency, gastrulation, differentiation, and organogenesis [5–7].

Sox4, Sox11, and Sox12 form the SoxC protein family. It has been shown that members of the SoxC family reveal overlapping expression and are functionally redundant [7–9]. In some tissues however, they differ in expression levels and transactivation rates [9]. During mouse development, SoxC genes show a widespread, largely overlapping expression pattern with highest levels in postmitotic neuronal progenitor cells of the neural tube, the dorsal root ganglia, the thalamus, the retina, and the cerebral and cerebellar cortex. In addition, transcripts are found in undifferentiated mesenchymal cells, the genital tubercle, endocardial cushions of the heart, the lung, the gut, the pancreas, and the nephrogenic mesenchyme [7]. The knock-out of either Sox4 or Sox11 in mice is lethal at E14 or directly after birth, respectively, due to severe
cardiac defects such as outflow tract malformations. In addition, these mice display further developmental defects including abnormalities in lymphocyte development (Sox4), or in eye and bone development (Sox11) (reviewed in [7]). In contrast, Sox12 null mice are viable and show no obvious malformations [9].

The expression and function of soxC genes during *Xenopus laevis* embryogenesis has not been investigated so far. Here we describe for the first time the spatiotemporal expression profile of sox4 and sox11 during early *Xenopus* development in detail. Functional analyses using specific morpholino oligonucleotides (MOs) targeting either sox4 or sox11 demonstrated a requirement of both genes during eye development, in particular for the formation and larvalization of the retina. In Sox4 or Sox11-deleted eyes, cell apoptosis was significantly induced whereas cell proliferation was not affected. Our data indicate an important role for Sox4 and Sox11 during vertebrate visual system development.

**Results**

**Sox4 and sox11 are Expressed in the Developing Eye of *Xenopus laevis***

A search in the GenBank database revealed a full-length cDNA sequence of the *Xenopus laevis* sox4 gene (Acc. No. NM_001098441). Based on this sequence, we successfully cloned a *Xenopus laevis* sox4 cDNA. Sequencing of several independent clones consistently revealed one amino acid exchange at position 180 from asparagine to serine in comparison to the published sox4 sequence and thus was considered to represent a polymorphism.

**Cloning of soxC**

We first determined the spatiotemporal expression of the three soxC members during early *Xenopus* embryogenesis. As we did not detect any significant expression of sox12 in the eye, we subsequently focused on sox4 and sox11 during early *Xenopus* embryogenesis. For this purpose, we generated antisense RNA probes that bind to the open reading frame of either sox4 or sox11 endogenous mRNA. Open reading frames of sox4 and sox11 revealed a nucleotide sequence homology of only 57%. Thus, we considered the probes to be specific for either sox4 or sox11. For detailed tissue-specific expression, we also generated vibratome sections of stained embryos (Fig. 1).

In the early *Xenopus* embryo, sox4 and sox11 were detected in the animal pole region of the embryo (Fig. 1A-A'). During gastrulation, sox4 and sox11 mRNA molecules were strongly visualized in mesodermal cells (Fig. 1B, B', C-C') and at stage 15 in the anterior neural plate (Fig. 1D-D'). Sox4 was additionally detected in the cardiac progenitor cell population (CPCs, Fig. 1D) and sox11 in the placodal primordium (Fig. 1D'). During late neurulation and tailbud stages, both sox4 transcripts were expressed in different brain regions, the posterior neural tube, the eye vesicles, the migrating neural crest cells, and the CPCs (Fig. 1E-H, K, E'-H', K'). During tadpole stages, sox4 was expressed in the first and the second heart field lineages, the vitelline veins and the aortic arch arteries (Fig. S1). At stage 34, sections revealed a strong sox4 expression in the mesod- and pericardium, and more faintly in the pericardial roof, the endo- and myocardium of the closed heart tube (Fig. S1). Sections through the head region of tadpole *Xenopus* embryos demonstrated that sox4 transcripts were located in the retina and the cornea epithelium of the eye as well as in defined regions of the brain (Fig. 1L-M). At stage 41, sox4 was expressed in the ciliary marginal zone (CMZ) of the eye (Fig. 1N).

Sox11 expression in the retina was strongly detectable in the ganglion cell layer forming a gradient towards more outer retina layers (Fig. 1L-M'), which is different to sox4 expression (compare to Fig.1L,M). At stage 41, sox11 expression was found in the CMZ identical to sox4 (Fig.1N').

In summary, sox4 and sox11 are strongly expressed in the developing *Xenopus* eye with an overlapping expression pattern suggesting a role for both genes during early eye development.

**Sox4 and Sox11 are required for *Xenopus* Eye Development**

Since sox4 and sox11 are specifically expressed in the *Xenopus* eye, we aimed to study the function of both genes during *Xenopus* visual system development by performing loss of function experiments using an antisense morpholino oligonucleotide (MO) based approach. For this purpose, we designed a MO targeting the translation start site of the endogenous sox4 mRNA (Fig. S2A). To investigate Sox11 function, we used a previously described Sox11 MO [10]. For rescue experiments, full-length human SOX4 or *Xenopus* sox11 constructs were injected which are not targeted by Sox4 MO (Fig. S2B) or Sox11 MO [10], respectively.

To test whether Sox4 or Sox11 have an influence on eye development, we injected several doses of both MOs into *Xenopus* embryos, fixed and investigated them at stage 41 when the retinal pigmented epithelium (RPE) is visible. Intriguingly, loss of either gene led to a quite similar eye phenotype. Depletion of Sox4 or Sox11 resulted in strong eye defects including smaller and deformed eyes whereas the un.injected side as well as the Control MO injected embryos developed normally (Fig. 2). Additionally, the RPE was not completely developed. Vibratome sections revealed severely disorganized retinal lamination upon Sox4 or Sox11 deficiency what we investigated in more detail using specific retinal marker genes (see below). To examine the specificity of the used MOs, we co-injected one of the MOs together with human SOX4 or *Xenopus* sox11 RNAs as mentioned above. Indeed, human SOX4 was able to significantly restore the eye defect induced upon loss of either Sox4 or Sox11 (Fig. 2 and Fig. S3). In addition, sox11 RNA was also able to revert loss of Sox11 as well as Sox4 function. These experiments clearly demonstrate the specificity of the MOs used and suggest that both genes are functionally redundant during eye development. Injection of Sox4 and Sox11 MOs together using low MO doses however did not reveal a cooperative effect of both genes (data not shown).

Taken together, these data indicate a requirement of Sox4 and Sox11 during *Xenopus* eye development.

**Sox4 and Sox11 Affect Retinal Lamination**

Since sox4 and sox11 are specifically expressed in the developing retina (Fig. 1L-M, L', M') and the depletion of both genes lead to severe eye defects including disarranged retinal layers (Fig. 2), we raised the question whether the formation of the different retinal cell types was affected upon Sox4 or Sox11 down-regulation. To examine these different cell types, we performed whole mount in situ hybridization experiments (WISH) using embryos at stage 41 with mild and severe eye phenotypes (Fig. 2) and probes for well characterized retinal cell specific marker genes [11]. Subsequently, we performed vibratome sections of stained embryos. We used ars3 and rho [12] to determine photoreceptor cells, px6 [13] to visualize amacrine and ganglion cells, vsx1 [14] to detect bipolar cells, prox1 [15] to show horizontal cells, and pmfjfl [16] to stain for ganglion cells. Almost all analyzed marker genes were expressed in the mildly and severely affected eyes. Only pmfjfl was absent in the severely affected ones (Fig. 3). These data indicate that all retinal cell types were formed excluding ganglion cells.

Intriguingly, we observed a severe disorganization of the different retinal layers as indicated by ectopic and missing
expression domains of marker genes (Fig. 3, red arrowheads). Furthermore bipolar and amacrine cells are displaced into the inner or outer layers of the retina (red arrowheads).

To analyze when this phenotype becomes apparent during development, we furthermore examined the same set of marker genes at stage 36 when the retina cell types start to differentiate.
We observed a similar phenotype at this stage (data not shown). Next, we investigated the expression of the proneural genes *rax*, *six3*, *pax6*, *otx2*, *pax1*, and *neurod1* at stage 32 before retinal cells are differentiated [17]. At this earlier stage, we could not detect a change in gene expression indicating that Sox4 and Sox11 have no influence on generating neuronal progenitor cells in the retina (Fig. 4).

In summary, Sox4 or Sox11 depletion had an influence on the fate of ganglion cell type and resulted in severely disturbed retinal lamination.

**Figure 2. Sox4 or Sox11 loss of function results in an abnormal eye development.** Effect of Sox4 (A) and Sox11 (B) depletion on eye development. The injection of Sox4 or Sox11 MO leads to smaller, deformed, and severely deformed eyes (white arrows) as illustrated in a dose dependent manner. A Control MO injection had no effect on eye development. Detailed views demonstrated defects in the formation of the RPE (red arrows). Vibratome sections showed that Control MO injection had no influence on the formation of the retinal layers and the RPE as well whereas depletion of Sox4 or Sox11 resulted in an abnormal RPE and retinal layering (red arrows). Quantitative representations are given. The eye phenotype of Sox4 or Sox11 down regulation was significantly rescued by the co-injection of either SOX4 or sox11 RNA. GCL = ganglion cell layer, INL = inner nuclear cell layer, n = number of independent experiments, N = number of injected embryos analyzed, ONL = outer nuclear cell layer, RPE = retinal pigment epithelium. Error bars indicate standard error of the means (s.e.m.), * P<0.05, *** P<0.001.

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**Sox4 and Sox11 in Lens Development**

To investigate lens formation upon Sox4 or Sox11 depletion, we injected both MOs independently and examined the expression of the two lens marker genes *cyr1a1* and *foxe3* at stage 36 (Fig. 5). As expected from *sox4* expression pattern in the eye (Fig. 1), loss of Sox4 had no effect on the expression of both genes while Sox11 depletion had a mild effect on both lens markers.
sox4 and sox11 in Xenopus
Sox4 and Sox11 in Early Eye Field Induction and Eye Differentiation

Beside the strong effect on retinal lamination, loss of either Sox4 or Sox11 function also led to the formation of smaller eyes in comparison to the uninjected side or Control MO injected embryos (Fig. 2). A possible reason for this phenomenon could be a defect in eye induction or early eye differentiation. Thus, we investigated the expression of the pan-neural marker gene sox3 as well as rax and pax6 as marker genes for the eye field at stage 13 in Sox4 or Sox11 depleted embryos. Neither Sox4 nor Sox11 function was required for the expression of sox3 and rax (Fig. 6A). Sox11 depletion led to a mild down-regulation of pax6 in the eye field. Moreover, we examined rax, pax6 and sox3 expression at stage 23 and observed a reduction in gene expression in the eye region in some Sox4 or Sox11 depleted embryos (Fig. 6B). Since sox4 and sox11 are also expressed in brain and neural crest cells, we

Figure 3. Sox4 depletion interferes with retinal lamination. Unilateral injection of 40 ng Sox4 or 40 ng Sox11 MO had no effect on the primary formation of most retinal cell types as shown by the expression of specific marker genes in mild eye phenotypes at stage 41. Only ganglion cells disappeared upon depletion of Sox4 or Sox11 in the severe eye phenotype. Many retinal cells were displaced (red arrowheads). Especially photoreceptor cells are displaced into inner layers of the retina (red arrowheads). In addition, the RPE is affected (black arrowheads). The uninjected (uninj.) sides revealed normal retinal lamination. For each marker gene, several embryos of different independent experiments were analyzed and showed a similar phenotype. Scale bar indicates 100 μm.
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Figure 4. Proneural genes are not affected upon loss of Sox4 or Sox11. At stage 32, expression of proneural genes was not changed after loss of Sox4 or Sox11. Scale bar indicates 100 μm.
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tested the expression of *emx1* (marker gene for the forebrain), *en2* (marker gene for the isthmus), and *egr2* (marker gene for specific regions of the hindbrain and migrating neural crest cells). Whereas all three analyzed genes showed a mild reduction upon loss of *sox11* function, only *en2* was reduced upon *sox4* knock-down (Fig. 6B). In all cases however, the phenotype had a low penetrance.

Taken together, we conclude that neural induction is not affected and early differentiation of neural tissue is only mildly affected upon *Sox4* or *Sox11* suppression. Defects in early neural specification can therefore be excluded as the main reason for the severe late eye phenotype.

**Sox4 and Sox11 in Regulation of Cell Proliferation and Apoptosis**

To clarify whether the smaller eyes observed upon *Sox4* or *Sox11* depletion was due to a misregulation of cell proliferation or apoptosis, we performed corresponding assays at stage 23, 32, and 41 using *Sox4* or *Sox11* deficient embryos. At stage 23, we could not observe any change in cell proliferation or apoptosis upon *sox4* depletion compared to the uninjected side or control MO injected embryos (data not shown).

Next, we analyzed cell apoptosis using TUNEL staining on whole embryos (Fig. 7A,B) and a caspase 3/7 enzymatic activity assay using isolated heads (Fig. 7C). At stage 32, loss of *Sox4* or *Sox11* led to a significant increase in TUNEL positive cells in the developing eye region and an increased caspase 3/7 activity compared to uninjected or control MO injected embryo. This increase in apoptosis persisted until stage 41 (Fig. 7D). *Sox4* has been described to be an anti-apoptotic factor by inhibiting Tp53 activity in hepatocellular carcinomas [18]. We therefore hypothesized that *Sox* MO injection may lead to an increased cell apoptosis in the *Xenopus* eye by activating Tp53. To test this hypothesis, we injected *Sox4* or *Sox11* MO together with Tp53 MO and observed a significant decrease in cell apoptosis compared to *sox* MOs co-injected with Control MO (Fig. 7E,F).

To investigate cell proliferation, we made use of pH3 staining. *Sox4* or *Sox11* depletion did not significantly alter cell prolifer-
sox4 and sox11 in Xenopus

A) 
- rax
- pax6
- sox3

Control MO

Sox4 MO

Sox11 MO

B) 

% embryos with reduced marker gene expression

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<th>Control MO</th>
<th>Sox4 MO</th>
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<td>rax</td>
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<td>pax6</td>
<td>n.s.</td>
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<td>sox3</td>
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N= 126 112 75 120 110 92 115 109 77

C) 
- rax
- pax6
- sox3
- emx1
- en2
- egr2

Control MO

Sox4 MO

Sox11 MO

D) 

% embryos with reduced marker gene expression

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<td>egr2</td>
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N= 99 78 71 127 80 68 67 80 96 122 70 72 130 86 75 107 108 73

n.s. = not significant
* = significant
** = highly significant
Sox11 null mice die immediately after birth as a result of cardiac defects [27]. Moreover, these mice reveal problems during eye, spleen, lung, and bone development. The consequence of Sox11 depletion especially on murine eye development has been investigated in more detail [23]. Sox11 knock-out results in severe defects during the formation of the cornea epithelium and lens fibers. In addition, retinal folds develop and the lens stays in contact with the overlying cornea epithelium (lens stalk).

One possible reason for the observed eye phenotype upon suppression of Sox4 and Sox11 function in Xenopus could be a perturbation of early neural or eye field induction and differentiation. In particular Sox11 gain-of-function in Xenopus animal cap cells results in neural induction [19]. Of note, loss of neither Sox4 nor Sox11 resulted in deficits in neural induction in our hands. We also investigated expression of pax6 since Pax6 depletion in the mouse reveals a similar eye phenotype as loss of Sox4 in our study [28]. We detected a mild but significant change in expression of pax6 and rax. This is in line with the observation of Sox11 null mice that do not show a severe alteration of Pax6 expression. Interestingly, Sox11 is under the control of Pax6 [23] and it remains to be investigated whether this is also the case in Xenopus.

In addition, proneural genes in the retina were not influenced upon loss of Sox4 or Sox11 function whereas ganglion cells differentiation was affected. These results are consistent with the observation that Sox4 and Sox11 act downstream of proneural genes but upstream of neuronal differentiation genes in the chick spinal cord [29].

**Sox4 and Sox11 in Cell Proliferation and Apoptosis**

SoxC genes have been shown to be involved in cell proliferation and apoptosis [22,23,30–32]. During sympathetic nervous system development, depletion of SoxC proteins leads to a decrease in BrdU-positive cells [22]. In contrast, Sox4 and Sox11 double-deficient mice show only mild changes in cell proliferation compared to WT controls during spinal cord development [30]. In mice, Sox11 depleted lens epithelium reveal a reduction in cell proliferation at E10.5, whereas at E9 no obvious effect could be observed [23]. In contrast, we did not see any significant effect on proliferation in the Xenopus eye at different stages upon Sox4 or Sox11 depletion. Of note, these data do not exclude that in Xenopus Sox depletion might result in reduced cell proliferation at other developmental stages in the eye (or other organs).

The available results concerning Sox4 function in cell survival and apoptosis are contradicting. On the one hand, it has been shown that SoxC genes in general are required for cell survival of neural as well as mesenchymal progenitor cells through the Hippo pathway [31]. In line with this, Sox4 and Sox11 have been described as survival factors during the development of the spinal cord [30] and sympathetic nervous system [22]. Repression of tp53 activity was observed by gain of Sox4 function [18]. In contrast, Sox4 was described as a DNA damage sensor in lung carcinoma cells [32] promoting cell cycle arrest and apoptosis. In our study, we could show that loss of Sox4 or Sox11 function leads to an increase in cell apoptosis in the Xenopus eye at stages 32 and 41 and this might well contribute to the small eye phenotype. Moreover, this phenotype was restored by inhibiting Tp53 providing a potential mechanism in which Sox4 and Sox11

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**Discussion**

In this study, we showed that (I) the HMG box transcription factors sox4 and sox11 are expressed in the developing Xenopus eye, (II) depletion of either sox4 or sox11 lead to severe malformations of the eye characterized by a decrease in size and disturbed retinal lamination, (III) loss of sox4 or sox11 does not affect neural induction, (IV) Sox4 and Sox11 have no influence on the primary fate of most retinal cell types and (V) Sox4 or Sox11 down-regulation results in increased cellular apoptosis in the eye.

**Sox4 and sox11 Expression Across Species**

We here provide a detailed description of the tissue specific expression of sox4 and sox11 during Xenopus embryogenesis and thereby extend earlier finding by others. The first cloning and maternal expression of sox11 was described earlier by others [19–21]. The expression of Sox4, Sox11, and Sox12 is highly conserved in mouse and chicken [7,8,22], in particular in the neural tube, the brain, the retina, the heart, and the kidney. A detailed expression of Sox11 in the developing murine eye has been published [23]. Interestingly, the expression pattern of murine Sox11 is quite similar to that of sox4 as well as sox11 during Xenopus eye development. Similar to our observation of sox4 and sox11 expression in the Xenopus eye, murine Sox11 is expressed in the evaginating optic vesicle. Later in development, Sox11 is strongly expressed in the retina, the lens fibers and the surface ectoderm. The RPE shows only a weak Sox11 expression. In zebrafish, the two duplicated sox11 genes, sox11a and sox11b, are also expressed in the developing eye and brain [24]. In contrast, sox4a is strongly expressed in the hindbrain and sox4b prominently expressed in the pancreas [25]. A second prominent expression site of sox4 and sox11 is the developing Xenopus heart. Sox4 is expressed in particular in the forming outflow tract. This is in line with published data in the mouse since murine Sox4 is also expressed in this region.

**Sox4 and Sox11 Function during Early Embryogenesis**

In our study, we demonstrated that the tissue-specific depletion of either Sox4 or Sox11 in the anterior-neural tissue leads to severe malformations of the eye including smaller and deformed eyes with a disorganized retina lamination. These phenotypes fit to the specific sox4 and sox11 expression in the different Xenopus eye structures. Sox4 knock-out mice die at E14 because of severe cardiac defects [26]. Especially the formation of the outflow tract is compromised in these embryos resulting in circulatory failures. In addition, lymphocyte development is disturbed in these embryos. The formation of other organs including the eye was not investigated in this study. Since sox4 is also expressed in the developing heart of Xenopus, it is certainly worthwhile to examine Sox4 function during Xenopus cardiac development by mesoderm-specific Sox4 MO injections.
function as survival factors during *Xenopus laevis* eye development [18,22,30].

### Experimental Procedures

#### Xenopus Laevis Embryos

*Xenopus* embryos were generated and cultured according to general protocols and staged according to others [33]. All procedures were performed according to the German animal use and care law (Tierschutzgesetz) and approved by the German state administration Baden-Württemberg (Regierungspräsidium Tübingen).

#### Cloning and MO Injection

For loss of function experiments, morpholino oligonucleotides (MOs) were designed and ordered from GeneTools, LLC: Sox4 MO: 5'-ACCATTGCTGCTGCTGTTTAGCTAC-3'; Sox11 MO: 5'-TCTGCTCGCTGCACCATGGCTGTCA-3' [10]; Tp53 MO: 5'-CCATGCCGGTCTCAGAGGAAGGTTC-3' [34]. Sox4 MO is designed to bind both published sox4 RNAs (sox4a: BC073494; sox4b: BC170171). For control injections, the standard control MO from GeneTools was used. All MOs were solved in DEPC-H2O and stored in aliquots at -20°C. For all experiments, 40 ng of the Sox MOs were injected unilaterally into one dorso-animal blastomere to target anterior neural tissue [35]. To analyze the role of Sox in apoptosis, 10 ng of Tp53 MO was coinjected with Sox MOs. 0.5 ng synthetic *gfp* RNA was coinjected as a lineage tracer in all experiments (see Fig. S2C). The uninjected side served as an additional, internal control. For rescue experiments, we used the full length human SOX4 construct of ImaGenes (clone #: IRAKp969B12110D) subcloned into pCS2+ vector [36]. To test the efficiency of the Sox4 MO, we cloned the *Xenopus* sox4 MO binding site (xsox4-gfp) and the corresponding region of the human construct (hSOX4 MO-gfp) in front of and in frame with the *gfp* open reading frame in pCS2+. 1 ng RNA of either fusion construct was injected together with either Control MO or Sox4 MO (Fig. S2B). Sox11 rescue RNA was used as described before [10]. RNA concentrations injected were: 0.7 ng *sox11* and 0.5–1 ng SOX4.

#### Whole Mount In Situ Hybridization

To analyze the spatiotemporal expression of *sox4* during *Xenopus laevis* embryogenesis, a full-length *sox4* cDNA was amplified with Phusion DNA-Polymerase (Biometra) from cDNA of stage 25 *Xenopus* embryos and ligated into the pCS-B vector (Stratagene). Primers used were: Sox4_f: 5'-TGC CCG GGG TGA CTG TAC TGC-3'; Sox4_r: 5'-TCA GTA GGT AAA TAC CAG GTT-3'. A DIG-labeled antisense RNA probe was generated by linearizing with ClaI (NEB) and in vitro transcription with T7 (Roche). Cloning and transcription of sox11 probe has been described earlier [10]. *Xenopus* embryos were fixed at different developmental stages with formaldehyde and WMISH analyzes were performed indicate standard error of the means (s.e.m.).

**Figure 7. Loss of Sox4/11 function leads to cell apoptosis in the developing eye.** 

**A:** TUNEL staining of Sox4 or Sox11 depleted embryos at stage 32. The areas where apoptotic cells have been counted are highlighted by a dotted circles and increased cell apoptosis was labeled by arrows. **B:** Quantitative representation of the TUNEL staining shown in A. n=number of independent experiments. **** P<0.0001. **C:** Increased caspase 3/7 activity after loss of Sox4 or Sox11 at stage 32. Values represent relative light units (RLU) normalized to Control MO injected embryos. n=number of independent experiments. Error bars indicate standard error of the means (s.e.m.). **D:** Unilateral injection of 40 ng Sox4 or Sox11 MO led to increased caspases 3/7 activity while uninjected sides or Control MO injected embryos were not affected. Values represent relative light units (RLU) normalized to the uninjected side. n=number of independent experiments. Error bars indicate standard error of the means (s.e.m.). **E:** Increased cell apoptosis upon loss of Sox4 and Sox11 was rescued by tp53 inhibition. **F:** A quantitative representation of the results in E is given. n=number of independent experiments, N=number of injected embryos analyzed. **G:** Crosssections of Sox4 or Sox11-depleted embryos demonstrating mitotic cells (blue, pH3 staining). Black arrowheads point to the MO injected side. **H:** A quantitative representation of the data in F is given. Sox4 or Sox11 depletion had not significant (n.s.) effect on cell proliferation in the eye at stage 32. n=number of independent experiments. Error bars indicate standard error of the means (s.e.m.), n.s. not significant, * P≤0.05, **** P<0.0001. doi:10.1371/journal.pone.0069372.g007
according to well-established protocols [37]. For a more detailed analysis of gene expression, we performed sections of stained embryos with a thickness of 25 μm using a vibratome (Leica).

**PH3 and TUNEL Staining using Whole Xenopus Embryos**

To detect cell proliferation as well as apoptosis in whole embryos upon Sox4 or Sox11 depletion, we performed pH3 (phospho histone H3) and TUNEL (Terminal Deoxynucleotidyltransferase-mediated dUTP Nick End Labeling) stainings according to standard protocols [37–39]. To investigate cell proliferation at stage 32, we made use of pH3 staining on whole embryos and counted pH3 positive cells on 7 μm plastic sections through the eye regions. We counted pH3+ cells in the eye on every third section (to prevent double counting of individual cells) and normalized the number of positive cells to the area covered by the eye.

**Caspase Assay**

For the analysis of cell apoptosis at stage 32, Sox4, Sox11, or Control MO was bilaterally injected into both animal-dorsal blastomeres and the head region of embryos was dissected at stage 32. For the analysis of cell apoptosis at stage 41, the eyes of unilaterally injected embryos were dissected at stage 41 (injected as well as uninjected sides). Head regions or isolated eyes were homogenized in 70 μl PBS/Triton X-100 (0.05 M sodium phosphate, 0.9% saline, 0.1% Triton X-100, pH 7.4) and protein concentrations were determined by Bradford assay using BSA as standard. Caspase 3/7 assays were done using the Caspase 3/7 Glo Assay (Promega, Madison, WI, USA) as described [40].

**Supporting Information**

**Figure S1 Transverse sections of an embryo at stage 34.**

**A:** Sox4 is expressed in the second (violet arrowhead) and first (red arrowhead) heart field lineage. **B:** Sox4 is expressed in the vitelline veins. **C:** Sox4 is expressed in the second heart field (black arrowhead) and the migrating neural crest cells (white arrow). **D:** Sox4 is detectable in the forming outflow tract (black arrow) and the ventral aorta/aortic arch arteries (black arrowhead). **E:** Sox4 is expressed in the mesocardium (me; black arrowhead) and the pericardium (p), and in the pericardial roof (pr), endocardium (e), and myocardium (m).

**Figure S2 Sox4 MO is specific.** **A:** Sox4 MO binding sites of Xenopus (sox4) and the corresponding region of human SOX4 (hSOX4). **B:** Co-injection of xSOX4 MO-gfp with control MO had no influence on gfp glowing. Sox4 MO blocked the translation of gfp. The human SOX4 binding site is not targeted by the Sox4 MO. **C:** The correct injection of Sox4 MO was controlled by gfp RNA co-injection. Dotted lines indicate the midline of the embryo.

**Figure S3 The eye phenotype after loss of Sox4 or Sox11 can be restored by both SOX4 and sox11 RNA.**

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**Author Contributions**

Conceived and designed the experiments: SJK MK TH. Performed the experiments: WC AH SJK ST MM. Analyzed the data: SJK MK TH. Wrote the paper: SJK MK.

**References**

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