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# Targeted resequencing identifies *PTCH1* as a major contributor to ocular developmental anomalies and extends the *SOX2* regulatory network

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Ocular developmental anomalies (ODA) such as anophthalmia/microphthalmia (AM) or anterior segment dysgenesis (ASD) have an estimated combined prevalence of 3.7 in 10,000 births. Mutations in *SOX2* are the most frequent contributors to severe ODA, yet account for a minority of the genetic drivers. To identify novel ODA loci, we conducted targeted high-throughput sequencing of 407 candidate genes in an initial cohort of 22 sporadic ODA patients. Patched 1 (*PTCH1*), an inhibitor of sonic hedgehog (SHH) signaling, harbored an enrichment of rare heterozygous variants in comparison to either controls, or to the other candidate genes (four missense and one frameshift); targeted resequencing of *PTCH1* in a second cohort of 48 ODA patients identified two additional rare nonsynonymous changes. Using multiple transient models and a CRISPR/Cas9-generated mutant, we show physiologically relevant phenotypes altering SHH signaling and eye development upon abrogation of *ptch1* in zebrafish for which in vivo complementation assays using these models showed that all six patient missense mutations affect SHH signaling. Finally, through transcriptomic and ChIP analyses, we show that *SOX2* binds to an intronic domain of the *PTCH1* locus to regulate *PTCH1* expression, findings that were validated both in vitro and in vivo. Together, these results demonstrate that *PTCH1* mutations contribute to as much as 10% of ODA, identify the SHH signaling pathway as a novel effector of *SOX2* activity during human ocular development, and indicate that ODA is likely the result of overactive SHH signaling in humans harboring mutations in either *PTCH1* or *SOX2*.

[Supplemental material is available for this article.]

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Severe congenital ocular malformations, including the reduction of eye size and anterior chamber anomalies at birth, are rare, with an estimated combined prevalence of 3.7 in 10,000 births (Bermejo and Martinez-Frias 1998). Ocular developmental anomalies (ODA) such as anophthalmia/microphthalmia (AM) or anterior segment dysgenesis (ASD) are underscored by extensive genetic heterogeneity, phenotypic variability, and nonpenetrance.

Anophthalmia refers to the complete absence of ocular tissue in the orbit (true anophthalmia), or the absence of ocular tissue upon clinical examination (clinical anophthalmia). Microphthalmia corresponds to a globe with a total axial length that is at least two standard deviations below the mean for age (<19 mm in a one-year-old child, <21 mm in an adult) (Weiss et al. 1989; Verma and Fitzpatrick 2007). To date, more than 20 genes have been implicated in AM in humans. Mutations in *SOX2* are the most frequent contributors to severe ODA (Fantes et al. 2003), yet account for a minority (10%–15%) of the genetic drivers (Verma and Fitzpatrick 2007), while other genes are more rarely involved. In AM cases, the genetic basis, either monogenic or chromosomal, is identified in ~20%–40% of individuals who undergo genetic testing (Slavotinek 2011; Chassaing et al. 2014). ASD disorders encompass a wide variety of developmental conditions affecting the cornea, iris, and lens including corneal opacity, posterior embryotoxon, iris hypoplasia, corectopia or polycoria, and adhesions between the iris and cornea or lens and cornea (Reis and Semina 2011). Two clinically distinct ASD disorders have been recognized as separate entities based on unique combinations of diagnostic criteria: Axenfeld-Rieger anomaly (iris hypoplasia, posterior embryotoxon, iris hypoplasia, corectopia/polycoria, and/or irido-corneal adhesions) and Peters anomaly (corneal opacity, defects in the posterior layers of the cornea, and lenticulo-corneal and/or irido-corneal adhesions). Causative mutations are identified in ~40% of individuals with Axenfeld-Rieger anomaly, mainly in two genes, *PITX2* and *FOXC1* (Pasutto et al. 2015). Despite the fact that thirteen genes have been associated with Peters anomaly, the genetic basis of this ocular anomaly remains unknown in the majority of cases (Weh et al. 2014).

Classical genetics approaches were successful to discover the first ODA genes. Linkage analysis studies allowed the identification of only few genes (Ferda Percin et al. 2000; Pasutto et al. 2007). The reason is most of the pedigrees available are small; therefore, the chromosomal regions that segregate with the disease are usually numerous and not reaching genome-wide significance. Positional candidate gene approaches pursuant to the identification of chromosomal abnormalities in ODA cases, or candidate gene approaches related to phenotype have yielded some gene discovery results such as *SOX2* and *OTX2* (Fantes et al. 2003; Ragge et al. 2005a). However, they have similarly reached an impasse toward identifying novel causal or contributory loci. In past years, the development of next-generation sequencing technologies has allowed the identification of several new ODA genes, including *ALDH1A3* and *RARB* (Fares-Taie et al. 2013; Srour et al. 2013). To identify novel ODA loci, we harnessed the power of next-generation sequencing technologies and conducted targeted high-throughput sequencing of a subset of candidate genes with a potential role in eye development, in a cohort of 22 sporadic ODA patients.

## Results

### Targeted resequencing identifies a significantly enriched mutational burden in *PTCH1*

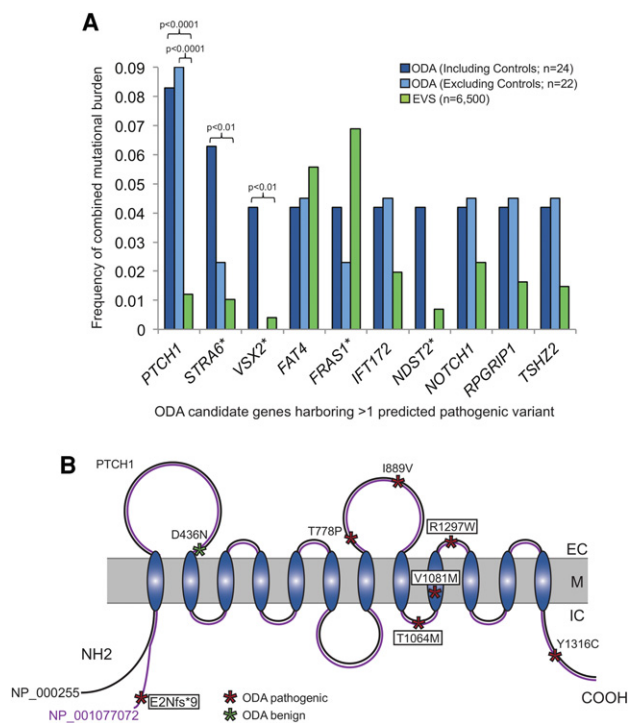
We selected 407 candidate genes (Supplemental Table 1) involved in ocular development for targeted exon liquid capture followed by

massively parallel sequencing. We screened 22 unrelated individuals with ODA and mutation negative for *SOX2*, *OTX2*, *RAX*, and *VSX2* and two positive control individuals (with known mutations in either *STRA6* or in *VSX2*). These affected individuals had isolated ASD ( $n = 6$ ), or AM that was isolated ( $n = 4$ ), associated with ASD ( $n = 6$ ), or coincident with coloboma ( $n = 6$ ) (Supplemental Table 2). We identified ~2500 variants in each patient; after stringent bioinformatic filtering that focused exclusively on alleles that were (1) absent from dbSNP132 (<http://www.ncbi.nlm.nih.gov/projects/SNP/>; Sherry et al. 2001), HapMap (<http://hapmap.ncbi.nlm.nih.gov/>; The International HapMap 3 Consortium et al. 2010), 1000 Genomes (<http://www.1000genomes.org/>; The 1000 Genomes Project Consortium 2015), and our in-house exomes; and (2) predicted in silico to be deleterious (Supplemental Table 3), we observed 0–5 variants per individual in a total of 46 loci. These genes harbored variation in the following locus-wide distribution: >1 variant/gene in 10 genes (Fig. 1A; Supplemental Tables 4, 5); 1 variant/gene in 36 genes; the remaining 361 genes were bereft of rare functional variants predicted to be deleterious (Supplemental Table 4, discussed in the Supplemental Note).

Among these 46 loci, *PTCH1* carried the greatest mutational burden and was the most significantly enriched for rare putative pathogenic variants in the 22 individuals from the ODA cohort and in the two positive control individuals C1 and C2 in comparison to 13,006 control chromosomes in the Exome Variant Server (EVS) ( $P < 0.0001$ ; Table 1; Fig. 1A,B) (<http://evs.gs.washington.edu/EVS/>; Fu et al. 2013) and remained nominally significant after correction for the 407 target gene set ( $P = 0.04$ ). Notably, the only other genes harboring an enrichment of rare variants predicted to be pathogenic in the 24 patients were *VSX2* and *STRA6*, both of which have been identified previously as rare ODA contributors (Ferda Percin et al. 2000; Chassaing et al. 2009) and were mutated in the positive control individuals C1 and C2, respectively ( $P < 0.01$  vs. 13,006 EVS chromosomes) (Fig. 1A; Supplemental Table 5). Upon exclusion of each of C1 and C2 from the mutational burden analysis, *PTCH1* was the only gene that carried a significant enrichment of pathogenic variants in the 22 ODA cases with previously unknown genetic etiology ( $P < 0.0001$  vs. EVS) (Fig. 1A).

Four individuals harbored rare *PTCH1* heterozygous changes predicted to be deleterious (Table 1; Fig. 1B). One patient with microphthalmia, cataract, and sclerocornea (P5) had a frameshifting deletion (c.4delG, p.Glu2Asnfs\*9) in exon 1 of *PTCH1* isoform NP\_001077072. Patient P20, affected with a bilateral Peters anomaly, harbored a heterozygous p.Tyr1316Cys change. Two additional unrelated patients with colobomatous microphthalmia, corpus callosum abnormality, and atrial septal defects (P8 and P15) harbored c.3191C>T (p.Thr1064Met) and c.3241G>A (p.Val1081Met) changes. With the exception of P5, for whom we were unable to perform segregation analysis, we determined that each of these three *PTCH1* mutations was inherited from an asymptomatic parent (Table 1; Supplemental Table 4), consistent with incomplete penetrance.

Because of the significant enrichment of *PTCH1* variants in our first-pass filtering strategy, and cognizant of the imperfect sensitivity and specificity of prediction algorithms, we returned to the set of rare variants remaining prior to in silico predictions. We found two additional heterozygous rare *PTCH1* missense variants filtered out initially because they were considered benign by PolyPhen-2 (<http://genetics.bwh.harvard.edu/pph2/>; Adzhubei et al. 2010). Individual P17, presenting with a bilateral Axenfeld-Rieger malformation, had a c.3889C>T (p.Arg1297Trp) mutation.



**Figure 1.** *PTCH1* has a significantly enriched mutational burden in ocular developmental anomalies (ODA). (A) Mutational burden for all genes harboring >1 rare predicted pathogenic variants in the initial cohort of ODA compared to healthy control individuals from the Exome Variant Server (EVS). Frequency of combined mutational burden is shown for the 10 genes harboring multiple rare (<1% alternate allele frequency) functional variants (frameshift, nonsense, and splicing variants were considered as damaging; missense variants were classified as damaging or not based on PolyPhen-2) in either ODA cases including two positive controls ( $n = 22$  unknown + 2 positive controls; dark blue bars), or ODA test cases alone ( $n = 22$  unknown; light blue bars) vs. EVS controls ( $n = 6500$ ; green bars).  $P$ -values are indicated for the only three genes with a significant enrichment in cases versus controls ( $\chi^2$  test). (\*) *VSX2* and *STRA6* are previously identified causal genes in the positive control ODA samples, C1 and C2, respectively; C1 also harbored variants in *FRAS1* and *NDST2*. (B) Schematic representation of the *PTCH1* receptor with its extracellular (EC), transmembrane (M), and intra-cellular domains (IC). The positions of the *PTCH1* mutations identified in ODA are represented with asterisks, with the color indicating variant pathogenicity. The four variants enclosed with boxes were identified in the initial discovery cohort. p.E2Nfs\*9 is specific to isoform NP\_001077072 (purple), which is identical to NP\_000255 (black) except for an alternate 66 amino acid region at the N terminus.

Additionally, we identified a heterozygous p.Asp436Asn change in the control sample C2 (Table 1; Fig. 1B).

### In vivo assay of variant pathogenicity shows that *PTCH1* variants are deleterious

The significant enrichment of mutational burden in the 24 cases compared to 6500 EVS controls ( $P < 0.001$ ), amino acid conservation, and in silico prediction evidence were suggestive but not conclusive with regard to the deleterious effect of the *PTCH1* missense variants identified in the ODA cohort (Table 1; Supplemental Fig. 1). Moreover, the observed incomplete penetrance posed interpretive challenges. Therefore, we evaluated the effect of all discovered alleles in vivo. *PTCH1* is a transmembrane-dependent receptor which functions with SHH as part of a dosage-sensitive pathway resulting in activation of downstream target genes, including the

Smoothed (*SMO*) coreceptor, *PTCH1* itself, and Gli transcription factors *GLI1*, *GLI2*, and *GLI3* (Villavicencio et al. 2000). SHH signaling is a key regulator of somite patterning (Bumcrot and McMahon 1995), and numerous zebrafish models of the SHH pathway including *shh* (Schauerte et al. 1998), *smo* (Varga et al. 2001), suppressor of fused (*sufu*) (Wolff et al. 2003; Koudijs et al. 2005), patched 2 (*ptch2*, referred to previously as *ptc1* [Koudijs et al. 2008]), and *kif7* (Tay et al. 2005) display a visibly more obtuse angle of the somitic chevron compared to wild-type (WT) controls. We have shown previously that in vivo complementation studies of human mutations in SHH effector molecules using somite defects as a phenotypic readout are a robust assay to determine allele pathogenicity (Putoux et al. 2011). We therefore employed this strategy to test the ability of human mRNAs harboring the ODA *PTCH1* missense mutations to rescue *ptch1* MO-induced somite angle defects in comparison to that of WT.

Using reciprocal BLAST, we confirmed that the single ortholog of human *PTCH1* is located on *Danio rerio* Chromosome 8 (73% identity, 80% similarity; human versus zebrafish protein); this gene, initially named *ptc2*, is mutated in the zebrafish *leprechaun* (*lep*) mutant (Koudijs et al. 2005) but will hereafter be referred to as *ptch1*. This locus is distinctly different from *ptc1*, the zebrafish ortholog of human *PTCH2* that is mutated in the zebrafish *blowout* (*blw*) mutant (Koudijs et al. 2008), and is now referred to as *ptch2*. We obtained a previously published splice-blocking (sb) morpholino antisense oligonucleotide (MO) targeting the exon 3 splice donor site (Koudijs et al. 2008), and we detected a partial suppression of *ptch1* expression by RT-PCR analysis of sb1 MO-injected whole-embryo lysates (Supplemental Fig. 2A). Even so, we observed somite phenotypes characteristic of Shh defects in WT embryos injected with 12 ng of *ptch1* sb1 MO (81.5° degrees vs. 97.5° degrees for control vs. *ptch1* sb1 MO;  $P < 0.0001$ ) (Fig. 2; Supplemental Table 7). Importantly, co-injection of 100 pg of capped human *PTCH1* WT mRNA resulted in a significant amelioration of the somite defect (85.9° vs. 97.5° for WT rescue vs. MO;  $P < 0.0001$ ).

We next evaluated the five missense mutations (p.Asp436Asn, p.The1064Met, p.Val1081Met, p.Arg1297Trp, p.Tyr1316Cys) identified among the 22 patients and two control ODA individuals. As a measure of the sensitivity and relevance of this assay, we also evaluated known deleterious variants that have been proposed to be pathogenic based on either genetic evidence or functional evaluation of SHH pathway readout. As such, we tested the effect of human *PTCH1* mRNA carrying the known pathogenic p.Thr1052Met change, associated previously with a holoprosencephaly (HPE)-like phenotype (including bilateral microphthalmia) and normal MRI (Ribeiro et al. 2006) or with alobar HPE (Ming et al. 2002). We also evaluated p.Leu360Arg, reported previously as a loss-of-function variant that is unable to complement *PTCH1* function in murine *Ptch1*<sup>-/-</sup> fibroblast cells (Bailey et al. 2003). Considering the possibility that the *PTCH1* variants in our ODA cohort could function as dominant negative alleles, we tested p.Gly509Val, a highly conserved change in the sterol-sensing domain that confers a dominant negative effect in *Drosophila* (Hime et al. 2004). As negative controls for the assay, we evaluated a nonsynonymous change found commonly in population controls, p.Pro1315Leu, and a second C-terminal change, p.Pro1125Leu, shown previously to restore SHH signaling readout in vitro (Bailey et al. 2003). In contrast to the significant rescue of the morphant phenotype resulting from co-injection of either WT mRNA or the negative control mRNAs (p.Pro1125Leu and p.Pro1315Leu), the morphant somite angle defect was

**Table 1.** *PTCHI* variants identified in ODA patients and/or studied in vivo using zebrafish experiments

Patient	Transcript ID	cDNA variation	Protein variation	ClinVar accession number	Inheritance	GERP score <sup>a</sup>	Grantham score <sup>b</sup>	Polyphen-2 Hum-Div	PolyPhen-2 Var	SIFT <sup>c</sup>	EVS	Protein location	Zebrafish studies
P5	ENST00000375274	c.4delG	p.Glu2Asnfs*9	SCV000259119	Unk	–	81	–	D(0.990)	–	Abs	–	–
P8	ENST00000331920	c.3191C>T	p.Thr1064Met	SCV000259128	Asy Fa	5.61	21	D(0.991)	P(0.782)	T(0.1)	2/13006	TM	LOF
P15	ENST00000331920	c.3241G>A	p.Val1081Met	SCV000259142	Asy Mo	5.32	101	B(0)	P(0.07)	D(0.02)	1/13006	EC	LOF
P17	ENST00000331920	c.3889C>T	p.Arg1297Trp	SCV000259148	Asy Fa	2.75	194	D(0.983)	P(0.541)	T(0.07)	1/12914	IC	LOF
P20	ENST00000331920	c.3947A>G	p.Tyr1316Cys	SCV000259154	Sy Mo	4.92	23	P(0.115)	P(0.066)	T(0.98)	9/12499	IC	LOF
C2	ENST00000331920	c.1306G>A	p.Asp436Asn	SCV000262553	–	5.63	38	D(0.999)	D(0.990)	T(0.18)	Abs	EC	Benign
CC-44	ENST00000331920	c.2332A>C	p.Thr778Pro	SCV000259171	Unk	5.73	29	B(0.03)	B(0.071)	T(0.5)	Abs	EC	LOF
CC-10	ENST00000331920	c.2695A>G	p.Ile899Val	SCV000259170	Asy Fa	3.49	81	D(0.951)	P(0.608)	D(0.02)	15/13006	EC	LOF
HPE <sup>d</sup>	ENST00000331920	c.3143C>T	p.Thr1052Met	–	Asy Fa	5.87	98	P(0.906)	B(0.444)	T(0.22)	3761/12568	IC	LOF
p.Pro1315Leu (rs357564)	ENST00000331920	c.3944C>T	p.Pro1315Leu	–	–	4.83	102	D(1)	D(0.994)	D(0)	Abs	EC	LOF
p.Leu360Arg <sup>e</sup>	ENST00000331920	c.1079T>G	p.Leu360Arg	–	–	6.07	97	D(1.00)	D(1.00)	D(0.02)	Abs	TM	DN
p.Gly509Val <sup>f</sup>	ENST00000331920	c.1526G>T	p.Gly509Val	–	–	5.34	94	D(1.00)	D(1.00)	D(0)	Abs	TM	Benign
p.Pro1125Leu <sup>e</sup>	ENST00000331920	c.3374C>T	p.Pro1125Leu	–	–	4.55					Abs	TM	Benign

(P, in first column) patient, (C) control, (CC) confirmation cohort, (HPE) holoprosencephaly, (LOF) loss of function, (DN) dominant negative, (Unk) unknown, (Fa) father, (Mo), mother, (Asy) asymptomatic, (Sy) symptomatic, (Abs) absent, (–) not available, (D) probably damaging, (P) possible damaging, (B) probably benign, (T) tolerated, (TM) transmembrane domain, (EC) extracellular domain, (IC) intra-cellular domain.

<sup>a</sup>The Genomic Evolutionary Rate Profiling (GERP) score ranges from –12.3 to 6.17, with 6.17 being the most conserved.

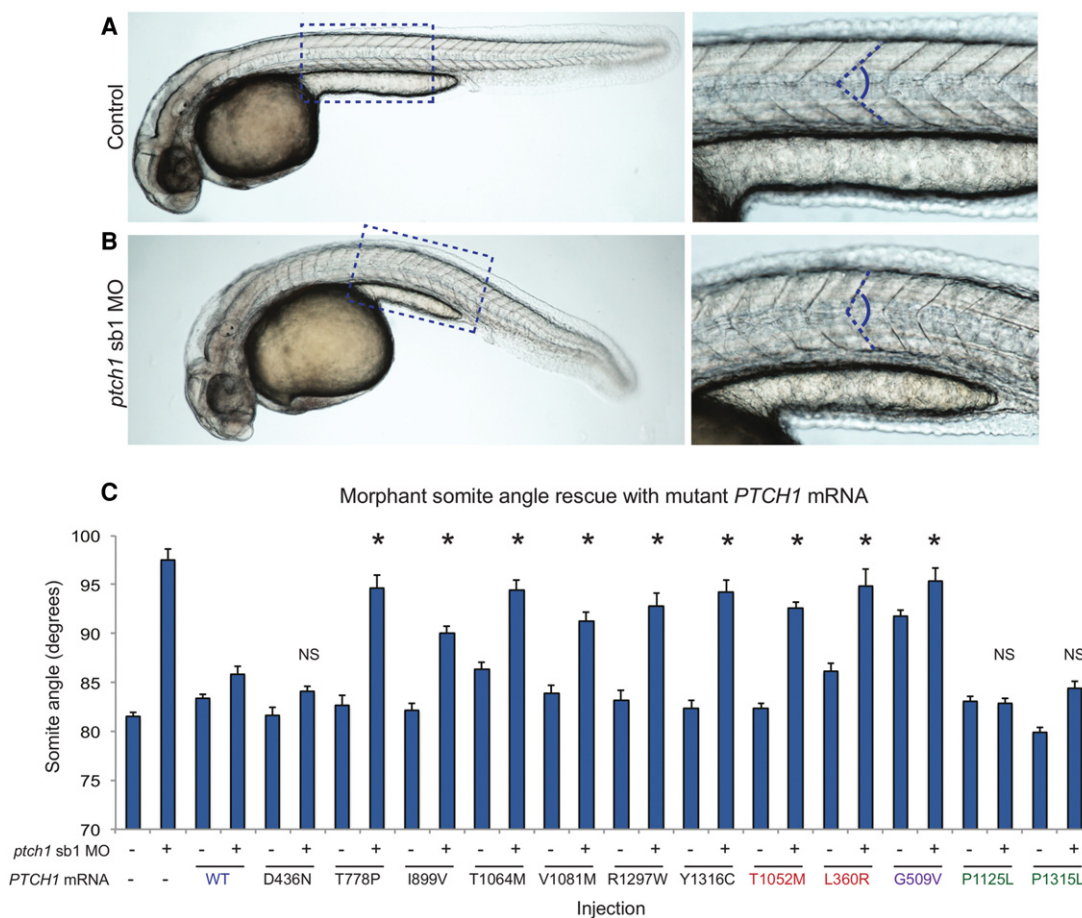
<sup>b</sup>Grantham scores, which categorize codon replacements into classes of increasing chemical dissimilarity, were designated conservative (0–50), moderately conservative (51–100), moderately radical (101–150), or radical ( $\geq 151$ ) (Li et al. 1984).

<sup>c</sup>SIFT (<http://sift.jcvi.org/>; Kumar et al. 2009).

<sup>d</sup>Ming et al. (2002).

<sup>e</sup>Bailey et al. (2003).

<sup>f</sup>Hime et al. (2004).



**Figure 2.** *PTCH1* variants identified in ODA patients are pathogenic. (A,B) Representative lateral images of uninjected control and *ptch1 sb1 MO*-injected live embryos taken at 36 h post-fertilization (hpf); dashed boxes are enlarged in the insets (right). Magnified panels show chevron-shaped somites (controls) and abnormal-shaped somites (morphants), caused by aberrant Hedgehog signaling in the zebrafish myotome. Dashed blue lines indicate measurement position (at the midpoint between the proximal hindgut and the anus) used for phenotypic scoring of embryo batches. (C) All six nonsynonymous *PTCH1* variants identified in ODA cases were pathogenic as indicated by the inability of mutant mRNA to rescue the *ptch1 MO*-induced somite angle defects. *PTCH1* p.Thr1052Met, a rare variant (minor allele frequency in controls 0.001;  $n = 13,006$  chromosomes [EVS]) reported previously in HPE, is also pathogenic; p.Leu360Arg is a previously reported functional null. Rescue with a common *PTCH1* p.Pro1315Leu encoding variant (rs357564; present in homozygosity in 8% of controls;  $n = 12,568$  chromosomes in EVS) was not significantly (NS) different from wild type (WT), nor was a previously reported benign variant p.Pro1125Leu change, providing support for the specificity of the assay. p.Gly509Val is a positive control for dominant negative effects. The missense variant p.Asp436Asn identified in the ODA control C2 was benign in this assay. We measured 38–58 embryos per injection batch with blind scoring. Asterisks indicate statistical differences between mutant and WT rescue ( $P < 0.0001$ ; Student's *t*-test). Error bars, SEM. See Supplemental Table 7 for somite measurement data.

significantly worse than WT rescue ( $P < 0.0001$  for each co-injection vs. WT rescue) and either partially ameliorated or not significantly different from *ptch1 sb1 MO* alone. We observed similar results for the p.Thr1052Met change (associated with HPE) and p.Leu360Arg (functional null) (Fig. 2; Supplemental Table 7). Notably, co-injection of *ptch1 sb1 MO* with the mRNA bearing the p.Asp436Asn change identified in control C2 showed phenotypic rescue similar to that of the WT and negative control mRNAs, demonstrating that this change is benign. Unlike the mRNA harboring the known dominant negative p.Gly509Val, injection of WT or mutant mRNA in the absence of MO resulted in no significant defects, arguing in favor of a loss-of-function rather than a dominant-negative effect for the ODA-associated alleles. Together, these results provided *in vivo* evidence that all rare missense variants identified in our ODA discovery cohort, as well as the mutation associated with HPE, have a detrimental effect on *PTCH1* protein activity (Supplemental Table 7), while the

negative control variants p.Pro1125Leu, p.Pro1315Leu, and the p.Asp436Asn change identified in a control patient had no detectable effect on protein function.

#### Pathogenic *PTCH1* alleles are present in an independent ODA cohort

Encouraged by these observations, we conducted bidirectional Sanger sequencing of the coding regions of *PTCH1* in an independent cohort of 48 samples with ODA. We identified two additional rare heterozygous *PTCH1* missense variants: p.Ile899Val in a patient with bilateral Peters anomaly, and p.Thr778Pro in an autosomal dominant AM-ASD family, each of which was absent from EVS (Table 1; Fig. 1B; Supplemental Table 4). *In vivo* functional assay of each of the two additional variants demonstrated that, similar to the alleles found in our original cohort, both changes resulted in partial loss of *PTCH1* function (mean somite angle

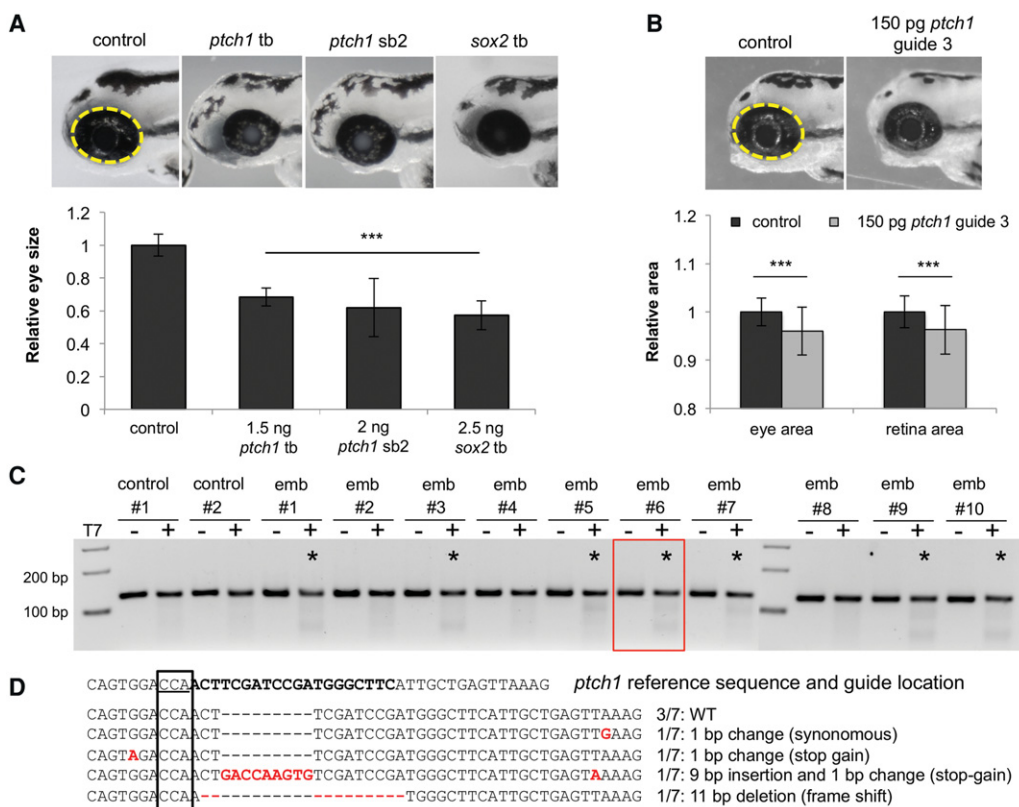
90.0° and 94.7° for p.Thr778Pro and p.Ile899Val, respectively;  $P < 0.0001$  for each MO plus mutant mRNA co-injection versus WT rescue (Fig. 2; Supplemental Table 6). Combined, we identified a total of seven rare heterozygous *PTCH1* variants (six missense and one frameshifting) in a total of 70 individuals with ODA (10%) (Fig. 1B).

### In vivo suppression of *ptch1* results in microphthalmia

Concomitant with aberrant somite formation, defects in the development of the visual system represent well-documented phenotypes in zebrafish SHH pathway component mutants. Importantly, zebrafish ENU mutant models of both patched homologs display ocular abnormalities: *ptch1* (*lep/ptc2*) mutants have vitreo-retinal abnormalities (Bibliowicz and Gross 2009); and mutation of *ptch2* (*blw/ptc1*) results in gross defects in eye morphology with variably penetrant coloboma (Karlstrom et al. 1996; Lee et al. 2008). To determine whether *ptch1* suppression produced a phenotype relevant to our ODA cohort, we evaluated *ptch1* sb1 MO-injected embryos for eye size defects by measuring eye area from lateral views at 3 d post-fertilization (dpf) and 5 dpf; consistent with previous observations, we observed no significant eye

size differences between sb1 morphants versus controls (data not shown; Koudijs et al. 2005). This is likely due to incomplete knock-down of endogenous transcript (Supplemental Fig. 2A). Therefore, we designed two additional MOs including a translation blocker (tb) and a splice blocker (sb2) targeting the exon 5 splice donor site of *ptch1*. RT-PCR monitoring of endogenous *ptch1* transcript in MO-injected embryos confirmed the higher efficiency of sb2 in suppressing WT *ptch1* expression in comparison to sb1 (Supplemental Fig. 2B). Next, we injected WT embryos with either *ptch1* sb2 or tb MOs; embryo batches were grown to 36 h post-fertilization (hpf) and scored first for abnormal somitic shape. Similar to *ptch1* sb1 MO-injected embryos, either mRNA splicing (sb2) or translational suppression (tb) resulted in obtuse somite angles significantly broader than controls (85.2° vs. 95.4° [tb] or 97.4° [sb2] for control vs. *ptch1* tb or sb2 MOs, respectively;  $P < 0.0001$ ) (Supplemental Fig. 3).

We then aged embryos to 3 dpf and the eye area was assessed as a percentage of WT eye area (Fig. 3A). We observed a significant reduction in eye size in both *ptch1* morphant batches (1.5 ng tb MO; 2 ng sb2 MO;  $n = 42$ –52 embryos/injection; repeated three times;  $P < 0.001$ ), supporting the direct role of *ptch1* in ocular development. Importantly, this phenotype was validated using a



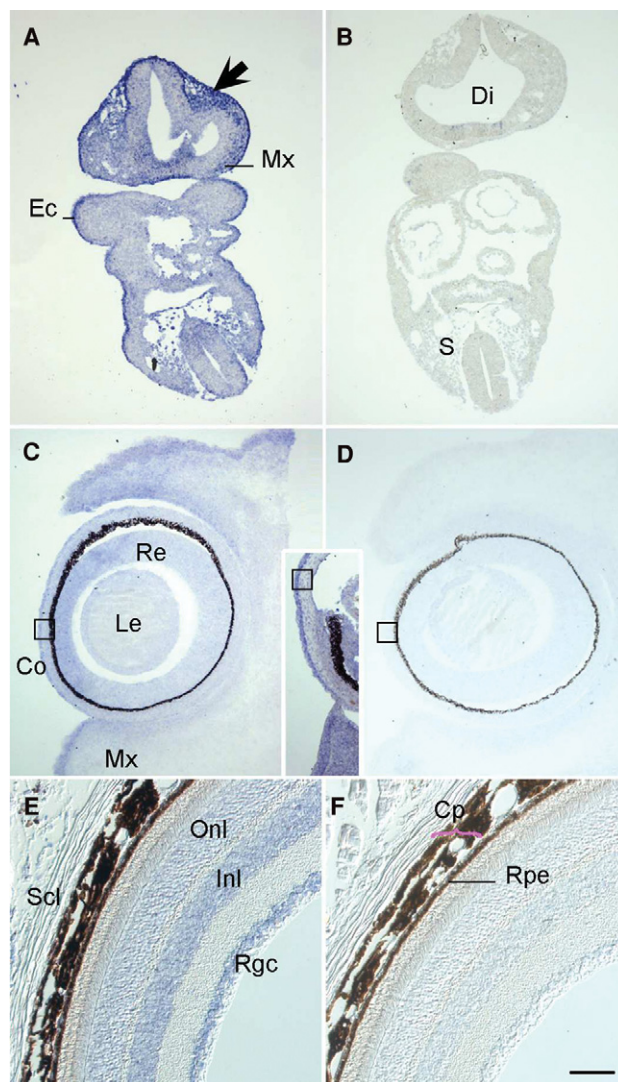
**Figure 3.** Suppression of *ptch1* in vivo results in microphthalmia. (A) Representative lateral images of uninjected control and *ptch1* tb, *ptch1* sb2, or *sox2* tb MO-injected embryos at 3 d post-fertilization (dpf) (top). Dashed yellow line indicates area of measurement used for quantification of eye size; morphant measurements were normalized relative to control (bottom). (B) Representative lateral images of uninjected control and *ptch1* guide RNA (gRNA)/Cas9 injected embryos at 3 dpf (top). *ptch1* CRISPR F<sub>0</sub> embryos were scored for eye size and retina size (bottom). (C) At 1 dpf, a representative sampling of 10 founders and two uninjected controls were selected and subjected to T7E1 endonuclease 1 (T7E1) assay. The appearance of T7E1 fragments < 100 bp (marked by asterisks) indicate positive gRNA targeting of exon 6 in the *ptch1* locus. No T7E1 fragments were detected in uninjected control embryos. Of the 10 founders subjected to T7E1 assay, seven showed the presence of T7E1 fragments, indicating that ~70% of founders have insertion/deletions (indels) in the exon 6 region of *ptch1*. (D) Multiple sequence alignment of *ptch1* reference sequence to *ptch1*-CRISPR variants generated from PCR amplification, subsequent TA cloning, and sequencing of *ptch1*-gRNA/Cas9 injected embryo #6. Black bold font marks the guide target, and the PAM recognition motif is underlined. Seven PCR-cloned sequences are shown, representing three wild-type variants and all four changes detected. TA cloning and sequencing of each founder embryo indicated 10%–65% mosaicism in individual fish ( $n = 5$  assessed).

CRISPR/Cas9  $F_0$  mutant model (Fig. 3B–D). Injection of a guide RNA targeting exon 6 of the *ptch1* locus induces gene-disruptive insertion-deletions (70% of embryos targeted; embryos with >65% mosaicism were selected for phenotyping) and leads to a significant reduction in eye size and retina size compared to controls ( $P < 0.001$ ) (Fig. 3B,C), as described for the *ptch1<sup>lep</sup>* mutant (Bibliowicz and Gross 2009).

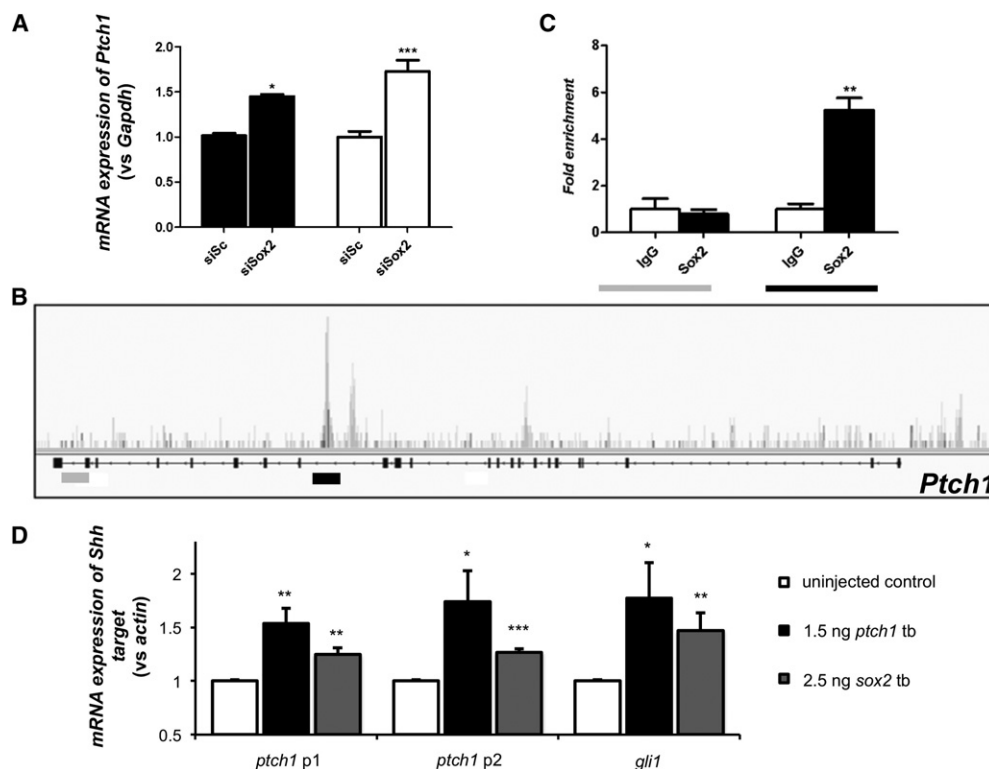
### *PTCH1* is regulated directly by the most frequently mutated ODA protein, SOX2

Pursuant to the elevated prevalence of ODA patients with pathogenic *PTCH1* variants, we wondered if this locus might be linked mechanistically to SOX2, as observed previously for other genes implicated in disorders of ocular development (Kamachi et al. 2001; Danno et al. 2008). We asked whether *PTCH1* might be regulated transcriptionally by SOX2. First, using RNA in situ hybridization, we found that robust embryonic *Ptch1* expression in the neural retina and lens persists to later stages in the adult mouse (Fig. 4), as observed in humans (Bakrania et al. 2008), and overlaps the known expression pattern of SOX2 (Hever et al. 2006). Using a physiologically relevant model of genetically modified murine stem cells overexpressing *Rax* (Tabata et al. 2004), we suppressed *Sox2* and tested the abundance of *Ptch1* message; in biological triplicate experiments, we found that *Ptch1* is up-regulated significantly upon *Sox2* suppression ( $P < 0.001$ ) (Fig. 5A), suggesting that the *Ptch1* locus might be under the transcriptional regulation of SOX2. We therefore performed chromatin immunoprecipitation (ChIP)-seq on CCE-Rx cells using an antibody against SOX2; we identified a peak in intron 15 of *Ptch1*, which was confirmed using targeted ChIP-qPCR on five independent samples. Importantly, amplification of the *Ptch1* intron 22 negative region was equivalent when precipitated with either nonspecific IgG or SOX2 antibody, while amplification of the intron 15 region showed greater than fivefold enrichment in chromatin immunoprecipitated by the SOX2-specific antibody ( $P < 0.01$ ) (Fig. 5B,C).

Next, we validated the putative regulation of *ptch1* by SOX2 in an in vivo context. First, we obtained a *sox2* tb-MO (Kamachi et al. 2008), and we evaluated eye size of zebrafish morphants. Consistent with both the ocular defects observed in humans with SOX2 mutations (Fantes et al. 2003; Chassaing et al. 2014), as well as the *ptch1* sb2 and tb-injected embryos, we observed a significant reduction in eye area upon translational suppression of *sox2* ( $P < 0.001$ ) (Fig. 3A). Further, we noted a significant defect in optic closure in *sox2* morphants (62% in morphants compared to 3.5% in controls,  $P < 0.001$ ), differing from *ptch1* morphant batches, which showed a modest proportion of embryos with coloboma (23.8% for *ptch1* sb2,  $P = 0.04$ ; 28% in *ptch1* tb  $P = 0.012$ ). Next, we monitored downstream SHH pathway activity in *sox2* morphants with qPCR analysis of *ptch1* and *gli1*. We observed significantly augmented *ptch1* expression in whole-embryo lysates from *sox2* morphants in comparison to controls at 2 dpf ( $P < 0.01$ ) (Fig. 5D). Notably, *ptch1* and *gli1* expression was also increased in *ptch1* tb-MO-injected embryos, suggesting that diminished PTCH1 protein results in overactivity of SHH signal transduction. These data are reminiscent of the transcriptional signature observed in vertebrate limb patterning in which SHH activation ensues after *PTCH1* inactivation (Butterfield et al. 2009). Since *PTCH1* is a regulator of its own expression through SHH pathway activity (Jeong and McMahon 2005), and SOX2 likely targets multiple SHH effectors (Zhao et al. 2012), deciphering the direct versus indirect roles of SOX2 protein on *PTCH1* expression remains challenging.



**Figure 4.** *Ptch1* transcripts are present in periorbital mesenchyme and the neural retina throughout eye morphogenesis and into post-natal life. (A) On embryonic day (E)9.5 in transverse mouse embryo sections, anti-sense riboprobes labeled with digoxigenin demonstrate *Ptch1* expression in cephalic mesectoderm of neural crest origin in the head (arrow) and maxillary arch (Mx); in the basal diencephalon (Di) and basal neural tube at trunk levels, and in the somitic sclerotome (S) and basal spinal cord. Ectodermal expression is constant at all embryonic stages examined (Ec). By E11.5, the distal diencephalic infundibulum transcribes *Ptch1* (data not shown) as does the subectodermal mesenchyme of the future eyelids and palate. (C) Mesenchymal *Ptch1* expression continues at E13.5, particularly in the superior and inferior palpebrae (eyelids; sPa, iPa); the lateral neural retina (Re) and differentiated outer cells of the lens (Le) begin to also transcribe *Ptch1*, which continues throughout these structures at E15.5 (inset). By this stage, initially generalized expression in the developing cornea has become restricted to the epithelium (Co, boxes in C/inset/D). (E) In adult mouse eyes on post-natal day 50, transcripts are found within the outer and inner nuclear layers (Onl, Inl), corresponding to photoreceptor and Müller cell bodies, and within the retinal ganglion cell layer (Rgc), testifying to a post-natal role in retinal maintenance. Transcripts not observed within the stroma of the anterior chamber or the sclera (Scl). (F) Rpe, retinal pigmented epithelium; Cp, choroid plexus. Scale bar A–D, 400  $\mu$ m; E, F, 200  $\mu$ m. Hybridization with a sense-oriented *Ptch1* probe as negative control in B, D, F.



**Figure 5.** SOX2 regulates *Ptch1* expression directly. (A) Targeted quantitative PCR after transfection of CCE-RX cells with a scrambled siRNA (siSc,  $n = 9$ ) or a siRNA targeting *Sox2* (siSox2,  $n = 9$ ). This experiment showed that decreased expression of *Sox2* leads to increased expression of *Ptch1*. (B) Results of ChIP-seq performed on CCE-RX cells using an antibody against SOX2. The *Ptch1* gene structure is represented underneath the DNA fragments sequenced after ChIP, with higher peaks corresponding to more enrichment. A peak was identified in intron 15 of *Ptch1* (underlined in black), while an example of an unenriched region is shown in intron 22 (underlined in gray). (C) Results obtained by ChIP-seq were confirmed using targeted ChIP-qPCR on five independent samples. Amplification of the intron 22 negative region (in gray) was equivalent whether using nonspecific IgG or a SOX2 antibody, while amplification of the intron 15 region (in black) showed greater than fivefold enrichment in chromatin immunoprecipitated by the SOX2-specific antibody. (A,C) Asterisks indicate statistical differences between the different conditions. (\*)  $P < 0.05$ , (\*\*)  $P < 0.01$ , (\*\*\*)  $P < 0.001$ , Mann-Whitney  $U$  test. Error bars, SEM. (D) Quantitative (q)PCR of *Shh* targets *ptch1* and *gli1* after *sox2* or *ptch1* tb morpholino injection in zebrafish embryos from three biological replicates. p1 and p2 indicate primer sets targeting two different regions of *ptch1*. (\*)  $P < 0.05$ , (\*\*)  $P < 0.01$ , (\*\*\*)  $P < 0.001$ , unpaired Student's  $t$ -test.

However, taken together, these data suggest that *PTCH1* expression can be regulated directly by SOX2 (as indicated by ChIP-seq) and that overactivity of SHH signaling conferred through the loss of function of either gene results in the same phenotypic consequences of ODA.

## Discussion

The SHH signaling pathway is associated strongly with ocular development in models ranging from insects to mammals that reflect ~600 million years of selection (Macdonald et al. 1995; Chiang et al. 1996; Nasrallah and Golden 2001; Takabatake et al. 2002; Bakrania et al. 2008; Christiansen et al. 2012). The *ptch1* zebrafish mutant displays an incompletely penetrant lens malformation phenotype (Koudijs et al. 2005); these mutants display abnormalities at the vitreo-retinal interface resulting in misshapen retinas and a reduced pupil size (Bibliowicz and Gross 2009). In addition, an ENU mutagenesis screen in zebrafish for visual system mutants identified a splice-acceptor site mutation in *ptch2* that results in ocular colobomas (Lee et al. 2013), and *ptch1;ptch2* double mutants have a severe ocular phenotype with absent lens development at 24 hpf and completely absent eyes at 48 hpf (Koudijs et al. 2008), phenocopying AM. Eye defects in *ptch1*<sup>ti222</sup> mutants could be suppressed by pharmacologically in-

hibiting the Hedgehog pathway with cyclopamine, providing evidence in support of a direct involvement of SHH signaling in the manifestation of the phenotype. Last, optic morphogenesis and gene expression patterns have been compared in blind cavefish and sighted surface fish embryos, both morphological variants of the same *Astyanax mexicanus* species (Yamamoto et al. 2004). In contrast to surface fish embryos, cavefish embryos develop small eye primordia, which later arrest in development, degenerate, and sink into the orbits, recapitulating human secondary anophthalmia. An expansion of the SHH signaling domain in the presumptive ocular neuroepithelium resulted in hyperactivation of downstream genes, lens apoptosis, and arrested eye growth and development in cavefish embryos (Yamamoto et al. 2004). These features could be mimicked in surface fish by *shh* overexpression, and eye development was restored partially in cavefish embryos by using cyclopamine (Yamamoto et al. 2004). Recently, it has been demonstrated that mutations in *SOX11* and *SMAD7* may lead to various ODA through the activation of the SHH pathway (Zhang et al. 2013; Pillai-Kastoori et al. 2014). In humans, *PTCH1* mutations have been associated previously with basal cell nevus syndrome (BCNS) (MIM#109400) and with holoprosencephaly (HPE7) (MIM#610828). Of note, ODA such as AM or ASD are part of both the BCNS (Bree and Shah 2011) and HPE (Pineda-Alvarez et al. 2011) phenotypes.

Our results show that the mutations identified in our ODA cohort lead to a loss of function of *PTCH1* and increased SHH signaling. We also show that inhibition of *SOX2* can drive overactivity of the SHH signaling pathway to produce the same ocular phenotypic outcome. Observations from our transient *ptch1* zebrafish models exemplify the dosage sensitivity of *PTCH1* in SHH signal transduction. The subeffective *sb1* results in normal eye size but a somite defect that can be rescued by co-injection with human *PTCH1* mRNA; however, the efficient *sb2* and *tb* MOs produced both a somite defect as well as a reduction in eye area in 3-dpf zebrafish embryos. Overexpression of human *PTCH1* mRNA in zebrafish embryos did not lead to an ocular phenotype (data not shown), suggesting that overexpression of this SHH target gene alone may not be sufficient to cause ODA. Moreover, although we demonstrated that *Sox2* regulates *ptch1* expression, it is probably not the primary mechanism governing eye development anomalies due to *SOX2* mutations, since *SOX2* is also involved in regulation of numerous genes including ocular developmental effectors such as *PAX6*, *OTX2*, and *RAX* (Hever et al. 2006; Danno et al. 2008).

Various ocular phenotypes (microphthalmia, colobomatous microphthalmia, and anterior segment dysgenesis) with incomplete penetrance are evident among our ODA cohort harboring *PTCH1* variants. Such a myriad of ocular phenotypes has already been associated with mutations in most of the genes involved in ODA (Slavotinek 2011), especially genes involved in the SHH pathway such as *SHH*, *GLI2*, and *PTCH1* (Roessler et al. 2003; Ragge et al. 2005b; Bakrania et al. 2010). The wide phenotypic spectrum within our ODA cohort, coupled to the incomplete penetrance observed among families, suggests that additional factors contribute to the phenotype. Such factors may include the other 45 genes harboring rare variants in the initial cohort of 22 ODA samples. However, their rarity in cases and undetectable enrichment of variation in comparison to controls suggests that their contribution will be modest; increased sample numbers and robust experimental models to test epistasis are required to demonstrate their involvement, if any, in ODA. Finally, phenotypic variability in ODA may also be explained by environmental factors, as suggested by the observation that mice heterozygous for a *Ptch1* null mutation had a fourfold higher risk to develop congenital malformations (including microphthalmia and anophthalmia) compared to WT mice when exposed to ionizing radiation during organogenesis (Hahn et al. 1998). More recently, mice heterozygous for a *Ptch1* null mutation were shown to develop spontaneous cataracts and increased susceptibility to cataract induction by early post-natal exposure to ionizing radiation compared to their WT counterparts (De Stefano et al. 2015).

In summary, high-throughput sequencing of candidate genes in ODA identified *PTCH1* as a significant contributor to congenital ocular malformations (10% in our cohort), placing it similar to its transcriptional regulator, *SOX2*, in terms of genetic burden to this phenotypic category (Fantès et al. 2003). Importantly, this study highlights the importance of a multifaceted approach toward identifying genetic contributors to traits such as ODA that are hallmarked by incomplete penetrance and genetic heterogeneity, especially when the cohort size is modest due to low disease frequency in the population. This study exemplifies how, together, a combined candidate gene sequencing approach, in vivo functional assessment of allele pathogenicity, and placement to a known disease gene network provides robust interpretive data that would not have been possible to achieve through genetic studies alone.

## Methods

### Candidate gene selection

We selected 407 candidate genes based on evidence for putative involvement in ODA (Supplemental Table 1). These genes were (1) linked to normal or abnormal ocular development in vertebrates and/or invertebrates in the literature, or (2) likely to be regulated by the *SOX2* transcription factor according to transcriptomic and ChIP-seq analysis.

### Patients

Seventy ODA patients (22 in the discovery cohort, 48 in the replication cohort) and two positive controls with known mutations were enrolled in this study. A signed informed consent was obtained from each participant, which adhered to the tenets of the Declaration of Helsinki and was approved by the local Ethics Committee (CPP Sud-Ouest and Outre-Mer II). Ocular phenotypes of the 22 patients included in the first cohort were isolated ASD ( $n=6$ ), isolated AM ( $n=4$ ), AM with ASD ( $n=6$ ), or AM with coloboma ( $n=6$ ) (reviewed in Supplemental Table 2). Ocular phenotypes of the 48 patients included in the second cohort were isolated ASD ( $n=14$ ), isolated AM ( $n=17$ ), AM with ASD ( $n=9$ ), or AM with coloboma ( $n=8$ ).

### Targeted enrichment and high-throughput DNA sequencing

A custom-made SureSelect oligonucleotide probe library was designed to capture the exons of 407 candidate genes (Supplemental Table 1). The probe library also aimed to capture 880 kb of potential regulatory sequences (i.e., noncoding region located within 20 kb of the 407 genes and conserved among species). A total of 56,059 probes, covering 2.46 Mb, were designed and synthesized. Sequence capture, enrichment, and elution were performed according to the manufacturer's instructions (SureSelect, Agilent). Each eluted-enriched DNA sample was then sequenced on an Illumina GAIIX as paired-end 75-bp reads (Integrage). Sequence reads were aligned to the reference human genome (UCSC hg19) using commercially available software (CASAVA1.7, Illumina) and the ELANDv2 alignment algorithm. The mean coverage was 325 $\times$  with 96.9% of the targeted sequences over 10 $\times$  and 93.4% over 25 $\times$ .

### Filtering strategy

All variants reported were filtered to ensure an optimal prioritization of candidate mutations (Supplemental Table 3). We first filtered out variants that did not meet the quality criteria (array confidence < 0.3, sequence read depth < 10, and sequence base quality < 10). We then filtered out all variants present in the local, in-house exome sequencing database (200 exomes) as well as in dbSNP132, 1000 Genomes Project, and the HapMap Project databases. Only exonic and splice-site variants were retained; we removed synonymous variants and variants predicted to be benign by the PolyPhen-2 software. The presence of the final selected variants was confirmed by Sanger sequencing.

### Mutational burden analysis

We determined the mutational burden for the 10 genes identified in the targeted resequencing of ODA samples harboring multiple rare (<1% alternate allele frequency) functional variants (frameshift, nonsense, and splicing variants were considered as damaging; missense variants were classified as damaging or not based on PolyPhen-2) in ODA cases ( $n=22$  unknown + 2 positive

controls) vs. Exome Variant Server controls ( $n = 6500$ ). EVS was accessed in November 2013, and a  $\chi^2$  test was used for comparisons.

### PTCH1 molecular screening

To support further the involvement of *PTCH1* lesions in ODA, we screened this locus in a new cohort of 48 patients by direct bidirectional Sanger sequencing. Primers used to amplify the 23 coding exons and intron-exon splice junctions are listed in Supplemental Table 7.

### Zebrafish embryo microinjection and manipulation

We obtained a previously published (Koudijs et al. 2005) morpholino antisense oligonucleotide (MO; GeneTools) targeting the splice donor site of *ptch1* exon 3 (sb1) and designed two additional MOs to suppress endogenous *ptch1* expression; a translation blocker and a splice blocker targeting exon 5 (sb2). We also obtained a previously described *sox2* tb (Kamachi et al. 2008; Supplemental Table 6). One nanoliter of the indicated cocktail was injected into wild-type (WT) zebrafish embryos at the one- to four-cell stage ( $n = 38$ – $58$  embryos/injection, repeated at least twice for MO plus mRNA injections and seven times for *ptch1* sb1 alone; with masked scoring).

For somite evaluation, embryos were reared at 28.5°C and imaged live at 36 h post-fertilization. To generate human *PTCH1* WT and mutant mRNA, we first obtained a full-length open reading frame (ORF) construct (clone ID: 100016192; OpenBiosystems). We generated a stop codon and subsequently introduced additional nonsynonymous changes using site-directed mutagenesis (QuikChange; Agilent). We then transferred sequenced-confirmed *PTCH1* ORFs into the pCS2+ plasmid (LR clonase II; Life Technologies), linearized with NotI, and performed in vitro transcription with the SP6 mMessage mMachine kit (Ambion). For rescue experiments, 12 ng of MO and 100 pg of *PTCH1* mRNA were injected, respectively. Live embryo imaging of lateral views was conducted on a Nikon AZ100 microscope at 6 $\times$  magnification facilitated by NIS Elements software. Somite angle measurements were taken at the midpoint between the proximal hindgut and the anus and recorded from the resulting images using ImageJ software. For eye size measurements, embryos were fixed in 4% PFA overnight at 3 dpf, and lateral images were acquired on an SMZ745T microscope facilitated by NIS Elements software. The area of the retina and pupil was quantified using ImageJ software. Embryo batches were compared for statistical significance using Student's *t*-test.

### Generation of CRISPR/Cas9 *ptch1* mutants

*ptch1* gRNA targeting the antisense strand of *ptch1* exon 6 was produced by synthesizing and annealing two oligonucleotides, gRNA3 F: TAGGGAAGCCCATCGGATCGAAGT and gRNA3 R: AAACACTTCGATCCGATGGGCTTC. The annealed oligos were then ligated to a BsmBI-digested T7cas9sgRNA2 vector overnight at room temperature (NEB). Two microliters of the reaction were used for transformation. Prior to transcription, the gRNA vector was linearized with BamHI. gRNA was transcribed using the MEGAShortscript T7 kit (AM1354, Life Technologies) and purified using alcohol precipitation. A total of 150 pg of *ptch1* gRNA and 300pg of Cas9 protein (PNA Bio) were co-injected into single cells of one-cell stage embryos. For the T7 endonuclease I assay, genomic DNA was prepared from 1-dpf embryos by digestion in 5  $\mu$ g/mL proteinase K for 90 min at 65°C, followed by 15 min at 95°C. A short stretch of the genomic region (~150 bp) flanking the *ptch1* gRNA target site was PCR-amplified from the genomic DNA (Fwd: TAGTGGCAAACCCCATATTAC, Rev: CCTTGACCCACATCTGCTTT). The PCR amplicon was then denatured slowly

and reannealed to facilitate heteroduplex formation. The reannealed amplicon was then digested with 5 units of T7 endonuclease I (New England Biolabs) at 37°C for 90 min. The samples were resolved by electrophoresis and ethidium bromide staining in a 3.0% agarose gel. PCR products were cloned (TOPO-TA cloning, Life Technologies), and individual colonies were sequenced to determine insertion-deletion sites and to estimate the extent of mosaicism in F<sub>0</sub> founders.

### Chromatin immunoprecipitation (ChIP)

We performed quantitative ChIP-seq in murine stem cells genetically modified to overexpress *Rax* (retina and anterior neural fold homeobox) (CCE-Rx cells, a kind gift from S. Watanabe) (Tabata et al. 2004). These cells have the ability to differentiate into retinal ganglion cells and were cultured using the standard procedures (Tabata et al. 2004). Two  $\times 10^6$  resuspended CCE-Rx cells were cultured in LIF-free medium on 10-cm bacterial plates. Forty-eight hours later, CCE-Rx embryoid bodies were treated with formaldehyde for 10 min, chromatin was prepared, and ChIP was performed according to the Upstate (Millipore) protocol, using 10  $\mu$ g of anti-SOX2 antibody (sc-17320, Santa Cruz Biotechnology) or mouse IgG (PP54, Millipore) as a control. ChIP-seq libraries were prepared and sequenced using the standard Illumina protocol. Peaks were called with SeqMonk using the contig generator function. Inside intron 15 of *Ptch1*, a peak was identified in the SOX2-immunoprecipitated sample. The online JASPAR database (<http://jaspar.genereg.net/>; Mathelier et al. 2014) confirmed the presence of a putative SOX2 binding site within the peak sequence. Amplicons corresponding to the *Ptch1* intron 15 SOX2 ChIP-seq peak and to a region not predicted to bind SOX2 in intron 22 (see Supplemental Table 8 for primers) were selected for validation. These amplicons were tested in five independent samples immunoprecipitated either with SOX2 or mouse IgG antibodies.

### Quantitative RT-PCR

Quantitative PCR analysis was performed in CCE-Rx cells and zebrafish embryos to confirm *Ptch1* regulation by SOX2. CCE-Rx cells were cultured using the standard procedures (Tabata et al. 2004) and transfected as previously described (Ko et al. 2009) either with an siRNA targeting *Sox2* mRNA or with a scrambled siRNA (Stealth siRNA MSS277200 and Negative Control Medium GC, respectively, Invitrogen). Quantitative real-time PCR analyses on complementary DNA transcribed from total RNA showed that the remaining *Sox2* expression was 50% at 24 h post-transfection and 75% at 48 h post-transfection. Forty-eight hours after transfection of either the siSOX2 ( $n = 9$ ) or the siScramble ( $n = 9$ ), total RNA was isolated using the GenElute Mammalian Total RNA Miniprep kit (Sigma-Aldrich). Samples were used to analyze *Ptch1* expression levels in the siSOX2 samples compared to controls (Supplemental Table 8 for primers). Zebrafish embryos injected with 1.5 ng *ptch1* tb MO or 2.5 ng *sox2* tb MO were grown for 48 h before isolation of total RNA using TRIzol (Invitrogen) following the manufacturer's protocol. cDNA was synthesized from 1  $\mu$ g of total RNA using SuperScript III (Invitrogen). Real-time PCR was performed with Power SYBR Green PCR Master Mix on a 7900HT (Applied Biosystems) and analyzed with a Sequence Detection System software package version 2.3 (Applied Biosystems) (Supplemental Table 8 for primers).

### In situ hybridization

In situ hybridization was carried out according to standard protocols (Goodrich et al. 1996) on paraffin sections using a murine digoxigenin-labeled *Ptch1* riboprobe (Chotteau-Lelievre et al.

2006) synthesized in antisense and sense (negative control) orientations.

## Data access

The data from transcriptomic and ChIP-seq from this study have been submitted to the NCBI Gene Expression Omnibus (GEO; <http://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE74600. All variants identified during this study have been submitted to the ClinVar database ([www.ncbi.nlm.nih.gov/clinvar/](http://www.ncbi.nlm.nih.gov/clinvar/); Landrum et al. 2016), and ClinVar accession numbers are listed in Table 1 (*PTCH1* variants identified in ODA patients) and in Supplemental Table S4 (final prioritized variants in patients with ODA).

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