PITX2 deficiency and associated human disease: insights from the zebrafish model

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Abstract

The PITX2 (paired-like homeodomain 2) gene encodes a bicoid-like homeodomain transcription factor linked with several human disorders. The main associated congenital phenotype is Axenfeld-Rieger syndrome, type 1, an autosomal dominant condition characterized by variable defects in the anterior segment of the eye, an increased risk of glaucoma, craniofacial dysmorphism and dental and umbilical anomalies; in addition to this, one report implicated PITX2 in ring dermoid of the cornea and a few others described cardiac phenotypes. We report three novel PITX2 mutations—c.271C>T, p.(Arg91Trp); c.259T>C, p.(Phe87Leu); and c.356delA, p.(Gln119Argfs*36)—identified in independent families with typical Axenfeld-Rieger syndrome characteristics and some unusual features such as corneal guttata, Wolf-Parkinson-White syndrome, and hyperextensibility. To gain further insight into the diverse roles of PITX2/pitx2 in vertebrate development, we generated various genetic lesions in the pitx2 gene via TALEN-mediated genome editing. Affected homozygous zebrafish demonstrated congenital defects consistent with the range of PITX2-associated human phenotypes: abnormal development of the cornea, iris and iridocorneal angle; corneal dermoids; and craniofacial dysmorphism. In addition, via comparison of pitx2M64* and wild-type embryonic ocular transcriptomes we defined molecular changes associated with pitx2 deficiency, thereby implicating processes potentially underlying disease pathology. This analysis identified numerous affected factors including several members of the Wnt pathway and collagen types I and V gene families. These data further support the link between PITX2 and the WNT pathway and suggest a new role in regulation of collagen gene expression during development.
Introduction

Paired-like homeodomain 2 (PITX2 [MIM 601542]) is a bicoid-like homeodomain transcription factor first identified as the cause of Axenfeld-Rieger syndrome (ARS), type 1 [MIM 180500], an autosomal dominant developmental disorder characterized by the combination of specific ocular and systemic defects (1–3). The typical ocular feature, often referred to as Axenfeld-Rieger anomaly, includes posterior embryotoxon, iris hypoplasia, iris processes, and corectopia/polycoria and is associated with a >50% risk of developing glaucoma (2,4). Systemic features include maxillary hypoplasia, micro- and hypodontia, redundant peribulbar skin and more rarely umbilical hernia, omphalocoele, anal anomalies, hypospadias, Meckel’s diverticulum and other defects (2,3,5,6). Several reports also implicate PITX2 in heart phenotypes including atrial fibrillation (AF) (7–11). In addition, one paper describes a family with a rare phenotype of ring dermoid of the cornea [MIM180550] carrying a pathogenic variant in PITX2 (12).

PITX2 has multiple isoforms, with three main transcripts seen in mammalian species: PITX2-A, -B and -C (13,14). The isoforms are generated by alternative splicing and/or use of different promoters and thus have diverse N-terminal regions; however, all three have identical homeodomain and C-terminal sequences. Most PITX2 mutations affect the homeodomain and C-terminal regions (via deletions, splicing or missense alleles) and result in complete or partial loss of function due to their disruption of all known PITX2 isoforms (3,6,13). Evident null alleles (complete gene deletions) support PITX2 haploinsufficiency as a mechanism of ARS (1,3,6,15). Human development appears to be highly sensitive to PITX2 dosage, as mutant proteins that retain limited wild-type activities seem to associate with milder phenotypes (16–18); in addition, increased transactivation capacity observed in some PITX2 mutants indicate that excessive PITX2 activity may also be harmful to proper development (6,19,20).

The normal developmental pathways downstream of Pitx2 are largely unknown; consequently, the mechanisms by which PITX2 mutations result in the variable Axenfeld-Rieger phenotype have yet to be fully elucidated. Dickkopf 2 (Dkk2), an extracellular antagonist of canonical Wnt-β-catenin signaling, was identified as the first downstream target of Pitx2 involved in pericardial mesenchyme/anterior segment development (21). Pitx2 has also recently been shown to be required for normal Tαp2b expression within the neural crest (22). In non-ocular tissues, PITX2 activates transcription of distal-less homeobox 2 (Dlx2) within the dental epithelium during tooth development (23). When considering co-factors of Pitx2, the most relevant protein-protein interaction to date is that of Pitx2 and forkhead box transcription factor C1 (Foxc1), another key gene implicated in ARS, type 3 [MIM 602482] (24,25). PITX2 and FOXC1 are co-expressed in distinct pericardial mesenchyme cells during multiple time points in ocular development, and both proteins localize to a common subnuclear compartment (26). In addition, the two proteins were shown to physically interact through immunoprecipitation assays; this interaction was found to rely on the PITX2 homeodomain and a PITX2-binding domain within the C-terminal activation domain of FOXC1 (26).

The function of PITX2 is highly conserved during vertebrate development. Mice homozygous for Pitx2 null or hypomorphomic alleles are embryonic lethal and display ARS-related anomalies including arrest in tooth development, abnormal eye placement, absence of ocular muscles and abnormal anterior segment structures, among numerous other abnormalities (27–30). In addition, heterozygous Pitx2 mutant mice previously considered unaffected have recently been shown to manifest the ocular features of ARS (31). Vertebrate conservation of PITX2 function also extends to zebrafish where two isoforms, pitx2a and pitx2c (orthologs of PITX2A and PITX2C, respectively), have been identified. Human and zebrafish ocular structures show significant anatomical similarities while also displaying a few differences. With respect to anterior segment structures, zebrafish and human corneas both display a stratified multilayer epithelium, a stroma of orthogonally-arranged collagen fibrils, and a monolayer endothelium (32–34); the remaining human corneal layers, Bowman’s and Descemet’s membranes, may not be present in zebrafish according to one recent paper (35). Zebrafish show a strikingly similar normal intraocular pressure (IOP) range to mammalian species, including humans (34). However, unlike in the human eye where aqueous humor is produced by the ciliary body and drained through the trabecular meshwork and Schlemm’s canal that extend circumferentially around the base of the cornea, aqueous humor production and drainage in zebrafish are managed by specialized structures located in the iridocorneal angle. Aqueous humor is produced in the dorsal angle of the eye by secretory cells of the non-pigmented ciliary epithelium, while drainage occurs in the ventral angle through the canalicular drainage network (32,36). Like humans, anterior segment structures in zebrafish receive contributions from diverse embryonic lineages (surface ectoderm, head mesoderm, neural crest and neuroectoderm), with pitx2 showing conserved robust expression in the neural crest-derived pericardial mesenchyme. Normal expression of pitx2 in embryonic tissues is critical to their development as complete morpholino-mediated knockdown of pitx2 results in ocular, craniofacial, and other defects (37). Recent publications also demonstrate the presence of a gross developmental phenotype in a zebrafish mutant line (38), as well as mild ocular defects in zebrafish mutants with deletion of the upstream regulatory regions of pitx2 that were previously shown to be conserved with humans (14,39).

In this manuscript we describe three novel dominant PITX2 mutations identified in three independent ARS families and highlight the variability of the associated phenotypes, both in terms of the severity of classic features as well as the occurrence of additional sporadic systemic defects. To gain further insight into the diverse roles of PITX2/pitx2 in vertebrate development and the mechanisms of associated variable phenotypes, we established permanent pitx2-deficient zebrafish lines via transcription activator-like effector nucleases (TALEN)-mediated genome editing and present their characterization with a focus on ocular abnormalities observed in homozygous animals. In addition, by comparing pitx2+/− and wild-type embryonic eye transcriptomes, we identify molecular changes associated with pitx2 deficiency that potentially underlie disease pathology.

Results

Identification of novel PITX2 mutations in human patients

Screening of PITX2 and FOXC1 in patients with ARS identified three novel causative mutations in PITX2: c.271C>T, p.(Arg91Trp) (Family 1); c.259T>C, p.(Phe87Leu) (Family 2); and c.356delA, p.(Gln119Argfs*36) (Family 3) (Table 1, Fig. 1). The first two mutations affect the homeodomain region of the protein, and the last one results in premature truncation and deletion of a large
Table 1. Summary of clinical and genetic features observed in Axenfeld-Rieger syndrome patients in this study

| Patient | Eye | Dental | Umbilicus | Other features | Race/ethnicity | DNA effect | Protein effect | Functional prediction | Family hx |
|---------|-----|--------|-----------|---------------|---------------|------------|--------------|----------------------|-----------|
| 1A      | ARA, GL, aniridia, partialis, coloboma | DA | RU | MH, BN, ptosis | Caucasian (Serbia) | c.271C>T | p.(Arg91Trp) | Damaging by 5/5 (S, PP, MT, MA, F) | Yes |
| 1B      | ARA, corneal guttata | DA | Unk | MH, BN | Caucasian (Serbia) | c.271C>T | p.(Arg91Trp) | Damaging by 5/5 (S, PP, MT, MA, F) | Mother of 1A |
| 1C      | ARA, GL, CA | DA | Unk | MH | Caucasian (Serbia) | c.259T>C | p.(Phe87Leu) | Damaging by 5/5 (S, PP, MT, MA, F) | Grandmother of 1A |
| 2A      | ARA, GL | Unk | Unk | MH | Caucasian (Serbia) | c.259T>C | p.(Phe87Leu) | Damaging by 5/5 (S, PP, MT, MA, F) | Yes |
| 2B      | ARA, GL | Unk | RU | MH, BN, (MR, SZ) | Caucasian (Serbia) | c.356delA | p.(Gln119Argfs*36) | Premature truncation | Son of 2A |
| 3A      | ARA, GL | HD, MD | RU, UH | mitral valve prolapse, WPW, hyperextensibility | Caucasian (USA) | c.356delA | p.(Gln119Argfs*36) | Premature truncation | Yes |
| 3B      | ARA, GL | HD, MD | UH | | Caucasian (USA) | c.356delA | p.(Gln119Argfs*36) | Premature truncation | Son of 3A |

ARA, Axenfeld-Rieger anomaly (any combination of posterior embryotoxon, iris processes, iris hypoplasia, and pupillary anomalies); BN, broad nasal bridge; CA, cataract; DA, dental anomaly; GL, glaucoma; HD, hypodontia; MD, microdontia; MH, maxillary hypoplasia; MR, mental retardation; RU, redundant umbilical skin; SZ, Seizures; UH, umbilical hernia; Unk, unknown.

<sup>a</sup>Numbering is relative to reference sequence NM_153427.2 where +1 is the A of the ATG initiation codon.

<sup>b</sup>5 prediction algorithms [SIFT (S), PolyPhen2 (PP), MutationTaster (MT), MutationAssessor (MA), FATHMM (F)] accessed through the Variant Effect Predictor (http://useast.ensembl.org/Homo_sapiens/Tools/VEP).

<sup>c</sup>MR and SZ felt to be due to neonatal asphyxiation, not PITX2 mutation.

Portion of the C-terminal region; even though all three mutations are novel, their expected functional effects are consistent with previously reported disease-associated alleles (Fig. 2). All three alleles are absent in over 250 000 general population alleles (gnomAD Browser). All three alleles co-segregated with the ARS phenotype in affected family members described below; no unaffected family members were available for testing. No other coding variants were identified in PITX2. Sequencing of FOXC1 in all three probands identified two common [c.1139_1141dupGCC, p.(Gly380dup)] and c.1359_1361dupCGG, p.(Gly456dup) at 16 and 25% allele frequency in the general populations, respectively, and one rare [c.1476_1481delGCCGCC, p.(Ala494_Ala499del)], at 0.073% allele frequency in-frame indels. Co-segregation studies were performed for the rare variant; however, all three met criteria to be classified as benign by ACMG/AMP (American College of Medical Genetics and Genomics/Association for Molecular Pathology) guidelines (40).

Patient 1A is an 8-year-old Caucasian male from Serbia diagnosed with ARS. He has bilateral typical ocular anomalies consisting of posterior embryotoxon, iris processes, iris hypoplasia and corectopia along with glaucoma diagnosed at 6 years of age. His left eye is more severely affected with partial aniridia and iris coloboma. Non-ocular anomalies include redundant periumbilical skin, dental defects, maxillary hypoplasia, broad nasal bridge, and ptosis (Fig. 1). His mother (Patient 1B) has posterior embryotoxon, iris hypoplasia, corneal guttata, dental defects, maxillary hypoplasia and a broad nasal bridge; no umbilical anomalies were noted. The patient’s maternal grandmother (Patient 1C) has posterior embryotoxon, iris hypoplasia, glaucoma and cataract, as well as dental defects and maxillary hypoplasia; no umbilical anomalies are noted. All three affected individuals were found to carry the heterozygous missense mutation c.271C>T, p.(Arg91Trp) in PITX2. While this specific mutation has not been reported previously, the mutation is predicted to be damaging by 5/5 in silico prediction packages, and a similar mutation affecting the same codon, c.272G>C, p.(Arg91Pro), has been reported (1). Screening of FOXC1 identified the benign c.1359_1361dupCGG, p.(Gly456dup) allele in Patient 1A.

Patient 2A is a 46-year-old Caucasian male from Serbia with ARS. He has bilateral posterior embryotoxon, iris processes, glaucoma and maxillary hypoplasia; dental and umbilical findings are unknown. His son, Patient 2B, also has bilateral posterior embryotoxon, iris processes and glaucoma. Non-ocular anomalies include redundant periumbilical skin, maxillary hypoplasia with a broad nasal bridge, and severe mental retardation and seizures thought to be caused by neonatal asphyxia. No details are available regarding his dental phenotype. Both Patient 2A and his son carry the PITX2 heterozygous missense mutation c.259T>C, p.(Phe87Leu), similarly predicted to be damaging by 5/5 in silico prediction packages. Screening of FOXC1 identified both common duplications listed above and the rare in-frame deletion; co-segregation analysis was performed for the rare benign variant and showed it was present in both affected family members.

Patient 3A is a 54-year-old Caucasian female from the United States with ARS. She displayed bilateral posterior embryotoxon, iris processes, iris hypoplasia, as well as glaucoma diagnosed at 25 years of age. Non-ocular anomalies include microdontia and hypodontia with 16 teeth missing (maxillary nos. 1, 2, 6-11, 14, 16; mandibular nos. 17, 20, 24, 25, 29, 32), redundant umbilical skin, umbilical hernia, and mitral valve prolapsa; no umbilical anomalies are noted. All three affected
prolapse. Her son (Patient 3B) is similarly affected with typical ARS ocular features and glaucoma diagnosed at 7 years of age, hypodontia, microdontia, umbilical hernia, Wolf-Parkinson-White syndrome (WPW), and hyperextensibility. Both patients were found to carry the heterozygous frameshift mutation c.356delA, p.(Gln119Argfs*36) in \( \text{PITX2} \). Screening of \( \text{FOXC1} \) identified the benign c.1139_1141dupGCG, p.(Gly380dup) allele in Patient 3A.

Generation of an allelic series of \( \text{pitx2} \) mutations in zebrafish

TALEN-mediated genome editing generated a series of mutations in the region encoding the second \( \alpha \)-helix of the \( \text{pitx2} \) homeodomain (Fig. 2). Sequencing of embryos produced by mosaic fish identified five mutant alleles including one premature truncation—c.190_197delATGTCGAC, p.(Met64*); c.193_195delTCG, p.(Ser65del); c.191_196delTGTCGA, p.(Met64_Ser65del); and c.193_195dupTCG, p.(Ser65dup)—all produced ~25% of embryos with defects in ocular and craniofacial development. Homozygosity for the corresponding \( \text{pitx2} \) mutations in all abnormal embryos was further confirmed by genotyping; no gross phenotype was detected in heterozygous embryos. Interestingly, the c.191_199delTGTCGA, p.(Met64_Thr66del) line produced a milder, incompletely penetrant phenotype with only 8% of homozygous embryos (2/24 in two distinct crosses) displaying the typical phenotype described above, indicating that this may be a partial loss-of-function (hypomorphic) allele.

The c.190_197delATGTCGAC, p.(Met64*) allele resulting in a premature stop codon and predicted to encode a truncated \( \text{pitx2} \) protein lacking the C-terminus and part of the homeodomain region. Embryos carrying each of the mutant alleles were raised to adulthood, genotyped and crossed to generate homozygous embryos for observation. Four mutant alleles—c.190_197delATGTCGAC, p.(Met64*); c.193_195delTCG, p.(Ser65del); c.191_196delTGTCGA, p.(Met64_Ser65del); and c.193_195dupTCG, p.(Ser65dup)—all produced ~25% of embryos with defects in ocular and craniofacial development. Homozygosity for the corresponding \( \text{pitx2} \) mutations in all abnormal embryos was further confirmed by genotyping; no gross phenotype was detected in heterozygous embryos. Interestingly, the c.191_199delTGTCGA, p.(Met64_Thr66del) line produced a milder, incompletely penetrant phenotype with only 8% of homozygous embryos (2/24 in two distinct crosses) displaying the typical phenotype described above, indicating that this may be a partial loss-of-function (hypomorphic) allele.

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p.(Met66Ilefs*133) mutation affecting the same methionine residue within the homeodomain that was identified in ARS (41).

**Gross morphological examination of pitx2<sup>M64+</sup> embryonic and adult phenotypes**

Morphological examination revealed that as early as 3-dpf (days post fertilization) the most prominent mutant ocular phenotype was a severely underdeveloped anterior chamber appearing bilaterally in 100% of homozygous pitx2<sup>M64+</sup> embryos (Fig. 3C' and D' versus C and D). Also around this timepoint, ~96% (127 out of 132) of homozygous pitx2<sup>M64+</sup> embryos demonstrated ventral iris colobomas either unilaterally or bilaterally (Fig. 3A' and B' versus A and B). By 5-dpf, in addition to the persistence of the coloboma (Fig. 3E' and F' versus E and F) and underdeveloped anterior chambers, mutant lenses of homozygous pitx2<sup>M64+</sup> embryos appeared elongated in the anterior/posterior plane compared with their spherical wild-type counterparts (Fig. 3G' and H' versus G and H). Craniofacial defects ranging from mild to severe also often accompanied the ocular anomalies. Alcian blue staining of seven 5-dpf mutants and three wild-type siblings showed variable degrees of improper positioning, size and structure of the components of the first (mandibular) and second (hyoid) pharyngeal arches (Fig. 3I versus J–L, M versus N and P).

Homozygous pitx2<sup>M64+</sup> mutants demonstrated reduced survival to adulthood (42 of 404, ~10.5%). Gross ocular phenotypic presentation in those who survived to adulthood ranged from anophthalmia (Fig. 3R) and severe microphthalmia (Fig. 3S) to iris coloboma/hypoplasia (Fig. 3T) and improper globe positioning (Fig. 3U); in some fish, ocular features showed asymmetric presentation (Fig. 3V). Approximately 67% (28 of 42) of mutant adult fish displayed bilateral microphthalmia/anophthalmia (severe), ~21% (9 of 42) showed extremely asymmetric phenotypes with one eye being affected with microphthalmia/anophthalmia/complete globe disorganization and the other eye being grossly normal in size but often misshapen and/or partly opacified (moderate), and ~12% (5 of 42) presented with bilateral normal-sized eyes with subtle shape irregularities and/or opacities (mild). Accompanying craniofacial deformities were also often visible. Incidentally, among homozygous fish surviving to adulthood, only females were capable of successful breeding.

**Analysis of ocular phenotype in pitx2<sup>M64+</sup> mutants by histology and electron microscopy**

To characterize the anatomical changes in the pitx2<sup>M64+</sup> homozygous mutant eyes, histological analyses including electron microscopy were performed. These analyses included adult mutants (6) and wild-types (3), 1-mpf (months post-fertilization) mutants (3) and wild-types (3), 14-dpf mutants (3) and wild-types (3) and numerous fish ranging from 3–5 dpf, including 17 mutants and 5 wild-types at 3-dpf. With electron microscopy (performed in 3- and 14-dpf wild-type and mutant embryos), focus was directed primarily on tissues arising from the neural crest, namely the corneal stroma and endothelium, the iris stroma, and the dorsal and ventral iridocorneal angles; development of these structures in zebrafish has been previously described, and our analysis of wild-type specimens corresponded well with the published observations (32). At 3-dpf, early formation of the anterior chamber can be observed as peripheral separation between the cornea and the lens (Fig. 4A). When considering the central cornea, the 3-dpf wild-type eyes display the expected three layers of stratification: epithelium, stroma and endothelium (Fig. 4B–D); because the anterior chamber is early in its development and has not yet fully formed, the central corneal endothelium is found flush against the lens capsule and epithelium (Fig. 4A, B and D). The corneal epithelium is delineated into surface and subepithelial cells which are further distinguished by the presence of many dark-staining small inclusion bodies versus fewer larger inclusion bodies, respectively (Fig. 4B and C). The wild-type corneal stroma is of uniform thickness and on its basal side borders the corneal endothelium (Fig. 4D). In contrast, the 3-dpf pitx2<sup>M64+</sup> mutant does not display distinct lens and cornea separation (Fig. 4A). The pitx2<sup>M64+</sup> mutant corneal epithelium contains smaller inclusion bodies than wild-type (in both surface and subepithelial layers) and instead exhibits small vacuoles in the surface layer (Fig. 4B' and C; Supplementary Material, Fig. S2). To the best of our knowledge,
Figure 3. Gross characterization of the pitx2M64* mutant phenotype. (A–H) Wild-type (A–H) and pitx2M64* mutant (A’–H’) embryos. At 3-dpf, prominent features include ventral iris coloboma (white asterisk in B’ versus B) and bilateral severely underdeveloped ocular anterior chambers (arrowheads in C’, D’ versus C, D). By 5-dpf, in addition to the persistence of the coloboma (white asterisks in F’ versus F) and underdeveloped anterior chambers (arrowheads in G’, H’ versus G, H), mutant lenses of
the significance of the inclusion bodies in the corneal epithelium of zebrafish embryos has not yet been determined. The mutant corneal stroma appears thicker in some places and less uniform than its wild-type counterpart; however, the difference is not significant. In addition, the mutant appears to lack a defined corneal endothelium and lens capsule, as the corneal stroma sits directly adjacent to the lens epithelial cells (Fig. 4D').

In terms of the iridocorneal region, at 3-dpf the pericellular mesenchymal cells of wild-types have begun their differentiation into irido stroma, consisting of pigment cells—namely xanthophores and iridophores—as well as embryonic vasculature and non-pigment cells (Fig. 4E–H). The iris pigmented epithelium has developed from the retinal neuroepithelium and is producing characteristic black melanosomes (Fig. 4E–H). In contrast, the iridocorneal angles of 3-dpf pitx2M64* homozygous mutants exhibit numerous structural differences from their wild-type counterparts. The composition of the anterior chamber fluid-filled space in pitx2M64* eyes is marked by an excess of cellular material (Fig. 4E–H'). Possible sources of this material include aberrant lipid or protein secretion, cellular sloughing/ degradation or undifferentiated cells which could be attributed to the notable under-differentiation of the irido stroma. While the dorsal iris stroma presents some evidence of delayed pigment differentiation and organization (Fig. 4E–F'), the ventral angle displays a complete lack of iris stroma differentiation (Fig. 4G–H'). In addition, though ventral iris pigmentation is generally delayed compared with dorsal pigmentation, the pitx2M64* mutant ventral angle shows a pronounced absence of melanosomes in the most anterior portion of the iris pigmented epithelium at this timepoint (Fig. 4G–H').

Examination of 14-dpf wild-type and pitx2M64* mutant embryos reveals persistence of the defects observed at the earlier stage (Fig. 5A versus A'). With respect to the cornea, the 14-dpf wild-type cornea is clearly stratified into its three components: a three- to four-cell layer thick epithelium; a smooth, even-thickness stroma; and a uniform single layer endothelium (Fig. 5B–D). The corneal endothelium is distinctly separated from the lens capsule and epithelium by the anterior chamber space filled with a proteinous liquid (aqueous humor) (Fig. 5B). The most noteworthy observation upon inspection of the 14-dpf pitx2M64* mutant eyes is the absence of the anterior chamber space, which places the mutant cornea directly up against the lens capsule (Fig. 5A–D'). The 14-dpf pitx2M64* mutant corneas seems to have three layers; however, the innermost layer of subepithelium, stroma, and endothelium appear wavy and of fluctuating thickness (Fig. 5B–D'). The presumptive corneal endothelium in particular is contoured between the stroma and lens capsule; before additional staining (see below), it was difficult to say whether the observed layer is in fact a monolayer endothelium or the compressed remnants of the anterior chamber space (Fig. 5C–D').

At 14-dpf, the anterior chamber is more developed and manifests as a defined space between the lens and cornea with separation between the iris and lens and no occlusion of the iridocorneal angles (Fig. 5A, E and H–J). Iridocorneal angles of 14-dpf wild-type eyes show further differentiation of the iris stroma as well as the presumptive annular ligament (Fig. 5E, G, I and K). In the dorsal angle, ciliary zone non-pigmented epithelial cells have begun to anteriorly differentiate into secretory cells (Fig. 5E and F). These cells are the source of aqueous humor production in zebrafish and will continue to develop a more secretory appearance (32,36). In the ventral angle, differences in specialization instead form structures specific for aqueous humor drainage. A canalicular drainage network consisting of two main branches arises over the first month of development; the iridocorneal canal presents as a channel within a non-pigmented mesothelial layer separating the annular ligament from the iris stroma, and the ciliary canal appears as an endothelium-lined passage through the pigmented epithelial base of the iris (32,36). In the ventral angle, formation of both channels of the canalicular drainage network can be observed by 14-dpf (Fig. 5I, K and L). Examination of the 14-dpf pitx2M64* mutant shows a notable reduction of the anterior chamber space between the cornea and the lens as well as at both iridocorneal angles, with the iris pigmented epithelium remaining markedly in contact with the lens (Fig. 5A', E', H', I' and J'). The iris stroma and presumptive annular ligament appear delayed in their differentiation (Fig. 5E', G', I', K' and L'). In addition, the dorsal non-pigmented ciliary epithelium does not appear to have begun differentiation, as the cells immediately adjacent to the iris pigmented epithelium maintain their retinal neuroepithelial morphology (Fig. 5E' and F'). In the ventral angle, the only indication of specialization is the break in the iris pigmented epithelium marking the presumptive location of the ciliary canal (Figure 5J' and K'); aside from this, the endothelial lining of the canal has failed to differentiate, and the iridocorneal drainage canal is undetectable (Fig. 5J', K' and L').

Histological sectioning of six mutant and three wild-type adult fish revealed a spectrum of gross phenotypes (Fig. 6A–C). The most prevalent phenotype includes the absence of both the anterior and posterior chambers as well as a severe reduction or complete absence of the iris (Fig. 6B and C). In the corneal region, drastic structural differences are observed in comparison to the wild-type counterpart. The adult wild-type corneal epithelium contains four to six layers of nonkeratinized, stratified squamous cells (33) as observed in our controls (Fig. 6D). In the mutant fish, the epithelium is thicker, and a subset of cells takes on a larger columnar morphology closely resembling that of the epidermal stratified epithelium surrounding the head (Fig. 6E and F). The corneal stroma of the wild-type is of uniform thickness, whereas the equivalent mutant layer is generally thinner, wavier in appearance, and sometimes interrupted (Fig. 6D and E). In place of the single-layer wild-type corneal endothelium, the analogous region in the mutant is a noticeably thickened, disorganized layer of cells of unknown origin (Fig. 6D–F). Aberrant pigmentation is also typically observed in the mutant corneal epithelium and endothelium (Fig. 6E–F). In addition, some of the eyes that appeared opaque or hidden by gross analysis were found to be covered by skin and cartilage structures growing together over the eye globe (Fig. 6B, C, E and F).
Electron microscopy of 3-dpf anterior chamber structures. (A, A’) Histological sections depicting orientation of wild-type (A) and pitx2
M64* mutant (A’) eyes. (B–D, B’–D’) Wild-type (B–D) and pitx2M64* mutant (B’–D’) central corneas. Wild-type eyes display three layers of stratification: epithelium, stroma and endothelium (B). The corneal epithelium is delineated into surface and subepithelial cells distinguished by many dark-staining small inclusion bodies versus fewer larger inclusion bodies, respectively (arrows in B, C). The wild-type corneal stroma is of uniform thickness and on its basal side borders the corneal endothelium (D). In contrast, corneal defects in the mutants include an epithelium that contains smaller inclusion bodies in both layers and exhibits small vacuoles in the surface layer (arrowheads in B’, C’), a thicker and uneven stroma (asterisk in D’), and lack of a defined endothelial layer (arrowhead in D’). Wild-type (E–H) and pitx2M64*, mutant (E’–H’) iridocorneal angles. Dorsal (E–F) and ventral (G–H) angles displayed early formation of the anterior chamber space as well as early differentiation of the iris stroma and iris pigmented epithelium. In mutants, both dorsal (E’, F’) and ventral (arrowheads in G’, H’) iridocorneal angles display notable under-differentiation of the iris stroma and iris pigmented epithelium, particularly ventrally; in addition, the anterior chamber space is marked by an excess of cellular material (asterisks in E’, H’). ce, corneal epithelium; cn, corneal endothelium; cs, corneal stroma; lc, lens capsule; le, lens epithelium; sce, corneal subepithelium; ac, anterior chamber; c, cornea; is, iris stroma; ipe, iris pigmented epithelium.
Figure 5. Electron microscopy of 14-dpf anterior chamber structures. Histological sections depicting orientation of wild-type (A) and pitx2<sup>M64*</sup> mutant (A’) eyes. Wild-type (B–D) and pitx2<sup>M64*</sup> mutant (B’–D’) central corneas. The wild-type cornea is clearly stratified into its three component layers: a three- to four-cell layer thick epithelium; a smooth, even-thickness stroma; and a uniform single layer endothelium (B–D). The corneal endothelium is distinctly separated from the lens capsule and epithelium by the anterior chamber space (B). In mutants, the anterior chamber space is absent, placing the mutant cornea directly up against the lens capsule (B’–D’). The innermost layer of subepithelium and stroma appear wavy and of fluctuating thickness (asterisks in C’, D’). The presumptive corneal endothelium, be it a monolayer endothelium or the compressed remnants of the anterior chamber space, is contoured between the stroma and lens capsule (arrowheads in C’, D’). Wild-type (E–L) and pitx2<sup>M64*</sup> mutant (E’–L’) iridocorneal angles. Wild-type dorsal (E–H) and ventral (I–L) angles display a distinct anterior chamber space with separation between lens and cornea and no angle occlusion (E, H–J); in addition, further differentiation of the iris stroma and the presumptive annular ligament is also demonstrated (E, G, I, K). Dorsally, the ciliary zone non-pigmented epithelial cells have begun to anteriorly differentiate from the retinal neuroepithelium (E, F). Ventrally, the formation of the iridocorneal canal and the ciliary canal, the two main branches of the specialized canalicular drainage network, can be observed (I, K, L). In contrast, the mutant shows a notable reduction in the anterior chamber space at both iridocorneal angles, with the iris pigmented epithelium remaining markedly in contact with the lens (asterisks in E’, H’–J’). The iris stroma and presumptive annular ligament appear delayed in their differentiation (E’, G’, I’, K’, L’). Dorsally, the non-pigmented ciliary epithelium does not appear to have begun differentiation (E’, F’). Ventrally, the only indication of specialization is the break in the iris pigmented epithelium marking the presumptive location of the ciliary canal (I’, K’); aside from this, the endothelial lining of the canal has failed to differentiate, and the iridocorneal drainage canal is undetectable (I’, K’, L’).
Taken together, this data suggests that complete deficiency in \textit{pitx2} results in abnormal development of the cornea, iris, and iridocorneal region and leads to severe ocular disorganization in adult fish.

**Analysis of \textit{pitx2}^M64* mutants with various markers**

To further examine the observed ocular phenotype in \textit{pitx2}^M64* homozygous mutants, additional analysis with various markers was performed. Periodic Acid Schiff staining (PAS) was used to visualize mucin-secreting goblet cells in the conjunctiva and other epithelium (21). N-cadherin (cdh2), which marks the corneal endothelium (29) as well as both the corneal and lens epithelia in zebrafish (42) was also employed. To visualize the corneal stroma, collagen1a1 (col1a1a), a member of the collagen 1 family which is a key component of stromal collagen fibrils, and corneal keratan sulfate proteoglycan (CKS) were utilized (33,49).
PAS staining demonstrated that in wild-type 1-mpf and older fish, mucin-secreting goblet cells are present in the eye periphery (Fig. 6G, H and J) but absent from the central corneal epithelium (Fig. 6G–I, K and L). However, goblet cells are detected throughout the entire corneal epithelium in age-matched pitx2M64* homozygous mutants, suggestive of a shift from corneal epithelial identity to an epidermis-like layer (Fig. 6M–Q).

Immunostaining of 1-mpf wild-type corneal stroma staining reveals a continuous, uniform stromal layer with col1a1a appearing both on the cornea (arrows in B and D) and around the eye (arrowhead in B) and CKS (A, C, D) being specific to the corneal stroma (arrows in C, D). In mutants, interruptions in or absence of col1a1a (E, F, H–J, L) and CKS (E, G–I, K, L) staining indicates disorganization of the corneal stroma (arrows in E–H, J–L). CKS also detects mucin-secreting goblet cells in the eye periphery and aberrantly approaching the corneal epithelium (I, K, L). Adult wild-type (M–O) and pitx2M64* mutant (Q–S) sections stained with col1a1a, CKS (red) and DAPI (blue). Wild-type adult corneal stroma continues to co-express col1a1a and CKS in a continuous layer (arrows in M and O) with CKS localized specifically to the cornea (arrows in N) and col1a1a also present around the cornea (arrows in M, N). In mutant adults, col1a1a and CKS staining is absent from the corneal stroma region (Q). Instead, col1a1a stains ectopic cartilage structures in the endothelial region and ocular periphery (asterisks in Q, K), whereas CKS stains mucin-secreting goblet cells in both the eye periphery and corneal epithelium (arrows in Q–S). Adult wild-type (P) and pitx2M64* mutant (T) sections stained with N-cadherin (green). In wild-type corneas, cdh2 detects both the corneal epithelium and the monolayer corneal endothelium (arrows in P). In mutants, while cdh2 staining was detectable but less robust in the epithelium, it did not appear in the endothelial region indicating the absence of a defined corneal endothelial layer (arrow in T). al, annular ligament; ce, corneal epithelium; cn, corneal endothelium; cs, corneal stroma; l, lens; re, retina.

Figure 7. Immunohistochemical analysis of 1-mpf and adult pitx2M64* mutant phenotype. 1-mpf wild-type (A–D) and pitx2M64* mutant (E–L) sections stained with col1a1a (green), CKS (red) and DAPI (blue). D, H and I represent magnified images of the corresponding boxed regions in A, F and I. Wild-type corneal stroma staining reveals a continuous, uniform stromal layer with col1a1a appearing both in the cornea (arrows in B and D) and around the eye (arrowhead in B) and CKS (A, C, D) being specific to the corneal stroma (arrows in C, D). In mutants, interruptions in or absence of col1a1a (E, F, H–J, L) and CKS (E, G–I, K, L) staining indicates disorganization of the corneal stroma (arrows in E–H, J–L). CKS also detects mucin-secreting goblet cells in the eye periphery and aberrantly approaching the corneal epithelium (I, K, L). Adult wild-type (M–O) and pitx2M64* mutant (Q–S) sections stained with col1a1a (green), CKS (red) and DAPI (blue). Wild-type adult corneal stroma continues to co-express col1a1a and CKS in a continuous layer (arrows in M and O) with CKS localized specifically to the cornea (arrows in N) and col1a1a also present around the cornea (arrows in M, N). In mutant adults, col1a1a and CKS staining is absent from the corneal stroma region (Q). Instead, col1a1a stains ectopic cartilage structures in the endothelial region and ocular periphery (asterisks in Q, K), whereas CKS stains mucin-secreting goblet cells in both the eye periphery and corneal epithelium (arrows in Q–S). Adult wild-type (P) and pitx2M64* mutant (T) sections stained with N-cadherin (green). In wild-type corneas, cdh2 detects both the corneal epithelium and the monolayer corneal endothelium (arrows in P). In mutants, while cdh2 staining was detectable but less robust in the epithelium, it did not appear in the endothelial region indicating the absence of a defined corneal endothelial layer (arrow in T).
Immunostaining with n-cadherin (cdh2) in wild-type corneas detected both the corneal epithelium and the monolayer corneal endothelium (Fig. 7P). In mutants, while cdh2 staining was detectable but less robust in the epithelium, it did not appear in the endothelial region, indicating the absence of a defined corneal endothelial layer (Fig. 7T). Thus, the unusual corneal region bears a striking morphological resemblance to the skin and connective tissue structures of the surrounding head, suggestive of a possible loss of corneal cell fate or a fate switch to epidermal skin at some point during development.

To identify the initial processes affected in pitx2 mutants that lead to the abnormal eye development, we analyzed migration patterns and cell death during the early steps of eye development. The migration of pitx2-positive pericellular mesenchyme was assessed using in situ hybridization (ISH) with pitx2- and foxc1a-specific probes in wild-type and mutant embryos. The abnormal pitx2 transcript expressed in homozygous pitx2M64* mutants was detectable at comparable to wild-type levels, suggesting that the mutant transcript is not subject to nonsense mediated decay. Staining with a probe recognizing both pitx2 isoforms performed in 48-hpf (hours post fertilization) embryos revealed no obvious differences in the distribution of abnormal pitx2 transcript in homozygous mutant embryos in comparison to normal pitx2 in wild-type fish. Staining for the foxc1a transcript, another marker of anterior segment development, similarly identified no obvious differences between pitx2M64* homozygous and control embryos (Supplementary Material, Fig. S1). This data suggests no obvious defects in migration of neural crest cells expressing mutant pitx2 to their proper locations within the developing eye.

Terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) staining to visualize possible aberrant increases in apoptotic cell death indicated that 48–120-hpf control embryos (with at least 10 embryos per stage) display few to no apoptotic cells in the cornea-lens region. The pitx2M64* homozygous mutant embryos at the same stages (48-hpf (16), 72-hpf (12), 96-hpf (5) and 120-hpf (8)) did not show a consistent increase in TUNEL staining overall but tended to have more apoptotic cells in the lens-cornea region, either as individual puncta or larger patches of stained cells at variable locations (Supplementary Material, Fig. S1); in addition to affecting only a subset of mutant fish, the observations were often unilateral. Though these experiments suggested an increase in apoptosis in some pitx2 mutants, the noted effects do not seem to be sufficient to explain the observed ocular abnormalities which, together with generally normal migration pattern, suggests that pitx2 function is required for proper differentiation of the cornea, iris and iridocorneal tissues.

Identification of downstream pathways affected in pitx2M64* embryonic tissues via global transcriptome analysis

To begin to ascertain the downstream pathways that are disrupted in pitx2M64* homozygous mutants, RNA from 23-hpf wild-type and pitx2M64* homozygous eyes only was submitted for microarray analysis. The pitx2M64* mutants show no visible phenotype at 23–24-hpf, with pitx2-positive pericellular mesenchymal cells surrounding the eye and completing their migration to the anterior segment region for subsequent differentiation. With this, the possible transcriptome changes detected at this developmental stage are expected to be due to deficiency in activating/repressing early pitx2 targets as well as their likely secondary effects rather than the result of the structural ocular abnormalities seen at later stages. The resulting list of differentially expressed genes identified 2226 down-regulated and 2044 up-regulated transcripts in pitx2M64* homozygous mutants as compared with wild-type controls. Full microarray data are available in the ArrayExpress database at EMBL-EBI (www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-6463.

The analysis was further focused on genes that demonstrated a significant decrease (2-fold or higher) in their expression, based on previous characterization of PITX2 as an activator (44). Pathway analysis of these significantly down-regulated transcripts using Ingenuity Pathway Analysis identified enrichment of several broad categories including cell movement, head development, and eye formation and morphology (Supplementary Material, Table S2), as well as signaling pathways, such as Wnt/j-catenin, PCP, calcium, actin cytoskeleton, Rho GTPase, epidermal growth factor (EGF), platelet-derived growth factor platelet-derived growth factor (PDGF) and integrin signaling. In several of these lists, the presence of multiple members of the WNT signaling pathway as well as collagen gene transcripts was notable. Analysis of the entire list of significantly (x2 or more) downregulated targets identified six WNT transcripts— wnt3, wnt4a, wnt5b, wnt7a, wnt9b and wnt10a, as well as multiple transcripts of tert encoding a transcriptional modulator of WNT/beta-catenin signaling (45), and seven collagen transcripts, col1a1a, col1a2, col5a2b, col5a3b, col7a1, col12a1 and col14a1, with orthology present in humans. Further analysis of these genes using publicly available resources (http://biogps.org; https://www.ncbi.nlm.nih.gov/pubmed/) revealed their expression in mouse ocular structures, with an enrichment in iris, cornea, eyecup and ciliary bodies similar to Pitx2 (Supplementary Material, Fig. S3), as well as possible roles in anterior segment phenotypes (46–52). Genes associated with the strongest expression/possible function in anterior segment structures, including all identified WNT transcripts and four collagens, col1a1a, col1a2, col5a2b and col5a3b, were selected for further investigation (Fig. 8 and Table 2). RNA transcript analysis by qPCR confirmed that all 11 selected targets were reduced at 24-hpf, as predicted by the microarray (Fig. 8). Apart from col5a2b and wnt7a, the other nine targets were also persistent in their down-regulation through 48- and 72-hpf, corresponding to the onset of the pitx2M64* homozygous phenotype (Fig. 8).

Discussion

Patients with PITX2 mutations usually display ARS with typical ocular abnormalities (posterior embryotoxon, iris hypoplasia and/or iris processes, pupillary anomalies) and other systemic features (most commonly dental and umbilical defects). More rarely reported ocular phenotypes include various corneal anomalies, such as Peters anomaly (characterized by corneal opacities, defects in the posterior layers of the cornea, iris processes and/or cornealirclear abnormalities), ring dermoid of the cornea (diagnosed based on the presence of limbal dermoids at the cornea/conjunctival border), or microcornea (a small corneal diameter) (2,3). Many ARS patients (~50%) develop an increased, and poorly manageable, IOP that leads to optic nerve damage and visual impairment; since the iridocorneal angle appears to be mostly open in ARS patients, the increased IOP is believed to...
be associated with incomplete formation of the trabecular lamellae and Schlemm’s canal (53).

The patients presented in this article demonstrated typical features of ARS along with some additional less frequent findings. One individual in Family 1 was reported to have corneal guttata; a literature search identified a few additional cases of ARS with corneal guttata/Fuch’s dystrophy with confirmed PITX2 mutations in two families (54–56). Corneal guttata affects the corneal endothelium and is part of Fuch’s dystrophy, which is characterized by the formation of lesions between the corneal endothelium and Descemet’s membrane resulting in death of the endothelial cells. Another patient in Family 3 was diagnosed with WPW. Some people with WPW display atrial fibrillation, which has been independently associated with alterations in PITX2 expression in patients without ARS (57). Other unusual features present in our patients included severe mental retardation and seizures observed in Patient 2B (though possibly related to neonatal asphyxiation) and mitral valve prolapse and hyperextensibility also seen in Family 3.

To gain insight into the mechanisms of PITX2-associated phenotypes, we generated mutations in zebrafish pitx2 that are expected to result in a complete knockout. The zebrafish pitx2M64* mutant line was selected for further studies. The zebrafish p.(Met64*) allele most closely resembles the human p.(Met66Ilefs*133) mutation that was identified in an ARS family with typical eye, craniofacial and dental features (no umbilical anomalies noted); one family member demonstrated a more severe ocular presentation with corneal edema, sclerocornea, and pigment deposition on the nasal and temporal peripheral corneal endothelium (41). The zebrafish pitx2M64* mutants exhibited related ocular and craniofacial defects. Ocular anomalies included all three layers of the cornea: absence of an identifiable corneal endothelium as early as 3-dpf and persisting through adulthood; disruption or absence of the stroma; and an apparent fate shift from corneal epithelium to head epidermis, complete with aberrant pigmentation and ectopic collagenous structures. In addition, from early embryonic stages the development of the iris, iridocorneal angles, and ciliary zone are significantly disrupted such that these structures are extremely reduced, lack proper differentiation, or are absent in most pitx2 mutant adult eyes. With respect to craniofacial development, pitx2 mutants display structural and positional malformations of the first and second pharyngeal arches.

Several findings observed in pitx2M64* fish deserve special attention. The first is the progressive loss of corneal identity, with the corneal epithelium taking on an epidermal-like fate. A similar phenotype has been previously reported in mice with temporal Pitx2 ablation that demonstrated a shift in gene/protein expression profiles of their developing corneas to more closely resemble conjunctival tissue (58). A suggested mechanism involves interaction with the WNT pathway. Mice with deficiency of Dkk2 (an extracellular antagonist of canonical Wnt-b-catenin signaling and a previously identified downstream target of Pitx2) show corneal abnormalities, including a complete fate shift from corneal epithelium to stratified epidermis in some animals; the shift is attributed to aberrant regulation of localized Wnt signaling throughout the ocular mesenchyme and limbus (21,59). In humans, one report connected a missense mutation in PITX2, p.(Arg62His), with ring dermoid of the cornea [MIM180550] in a large Chinese pedigree (12,60,61). Genetic studies in other human pedigrees affected with corneal dermoid and related phenotypes (62) may provide additional information about possible other PITX2 mutations in these conditions. Also, the data presented in this manuscript provide further support for the interaction between PITX2 and the WNT pathway and encourage further investigation into their relationship (see below).

Another relevant observation noted in pitx2M64* adult zebrafish is the formation of ectopic cartilaginous structures appearing to
from the limbus and extend into the presumptive cornea. It could be hypothesized that pitx2 within the ocular mesenchyme serves not only to maintain corneal identity but also to counteract aberrant cartilage formation, so that when depleted, the mesenchyme and surrounding cells shift their fate to skin and cartilage. An alternative hypothesis is that presumptive iris stroma or corneal endothelial cells are failing to properly differentiate but instead either become and/or maintain a multipotent stroma or corneal endothelial cell fate to remain behind the lens and leading to retinal detachment. Additional studies are needed to better understand the possible role of PITX2/pitx2 in the vitreous/posterior segment.}

A final notable feature of pitx2<sup>−/−</sup> animals is a reduced or absent posterior vitreous chamber that, together with the lack of anterior chamber, leads to an overall more compressed appearance to the eye. In mammals, the vitreous body is an extracellular matrix reservoir of basal membrane proteins derived from the ciliary body and lens (64, 65). The vitreous body of zebrafish should be developed by 1 mpf and is delineated from the anterior chamber by the formation of lens zonules (32). Though it has not been studied in zebrafish, the failure of proper ciliary zone differentiation in pitx2<sup>−/−</sup> mutants may account for the absence/reduction of the vitreous body due to the lack of secreted vitreous components. The most commonly noted posterior defect in ARS patients is glaucomatous cupping of the optic nerve head, which is believed to develop in response to the increased IOP. However, a literature search identified several reports that presented additional posterior segment anomalies in ARS. A ‘primary vitreous condensation’ was noted in a patient with bilateral microphthalmia, sclerocornea, and ptosis, small teeth, bifid uvula, hearing and other defects (46). Dyskeratosis congenita, AD/AR, 613989; Leukemia, acute myeloid, AD, 613989; Melanoma, cutaneous malignant, 9, 615134; Pulmonary fibrosis and/or bone marrow failure, 1, AD, 614742 appearance to the eye. In mammals, the vitreous body is an extracellular matrix reservoir of basal membrane proteins derived from the ciliary body and lens (64, 65). The vitreous body of zebrafish should be developed by 1 mpf and is delineated from the anterior chamber by the formation of lens zonules (32). Though it has not been studied in zebrafish, the failure of proper ciliary zone differentiation in pitx2<sup>−/−</sup> mutants may account for the absence/reduction of the vitreous body due to the lack of secreted vitreous components. The most commonly noted posterior defect in ARS patients is glaucomatous cupping of the optic nerve head, which is believed to develop in response to the increased IOP. However, a literature search identified several reports that presented additional posterior segment anomalies in ARS. A ‘primary vitreous condensation’ was noted in a patient with bilateral microphthalmia, sclerocornea, and ptosis, small teeth, bifid uvula, hearing and other defects (46). Dyskeratosis congenita, AD/AR, 613989; Leukemia, acute myeloid, AD, 613989; Melanoma, cutaneous malignant, 9, 615134; Pulmonary fibrosis and/or bone marrow failure, 1, AD, 614742...
These distinctions may be attributed to differences in PITX2/pitx2 dosage requirements between species. Homozygous mutations in human PITX2 have not been reported but are likely to result in more severe ocular malformations, possibly resembling the spectrum observed in pitx2<sup>2Mel</sup> homozygous animals. There is evidence to suggest that some PITX2-positive cases affected with a more severe phenotype may have a greater disruption of PITX2 function. For example, a more severe phenotype was identified in a patient with a dominant-negative mutation in PITX2 (70). In addition, patients carrying mutations in both PITX2 and FOXC1, shown to result in a cumulative effect with greater loss of function, displayed severe ASD with corneal opacification, peripheral corneal vascularization, and PHPV ultimately resulting in bilateral retinal detachment (71). Histological analysis of the affected cornea showed corneal epithelium focal thickening and basal vacuolation, stromal vascularization, stromal collagen fiber separation due to accumulation of mucopolysaccharide-like ground substance, and partially or fully absent Bowman’s layer, Descemet’s membrane, and endothelium (71). It is also interesting to note that, even though the visible abnormal phenotypes in zebrafish are present starting at embryonic stage 3 dpf, the features become more severe as animals age. Although ARS in human patients is often diagnosed at birth (with features like, iris hypoplasia/iris processes being notable at that point), some attributes (like increased IOP and glaucoma) are known to develop with age. It is possible that phenotypic progression in humans involves additional ocular features that have not yet been recognized due to the lack of longitudinal studies.

Transcriptome analyses to identify ocular factors that are dependent on normal pitx2 expression during early eye development revealed numerous transcripts, with several selected genes of WNT and collagen families verified by quantitative PCR. As discussed earlier, the link between the Wnt pathway and Pitx2 has been previously identified. Not only has the Wnt/Disheveled/β-catenin pathway been shown to both directly induce Pitx2 expression and stabilize Pitx2 mRNA in a cell-type-specific manner, but Pitx2 has also been demonstrated to regulate Wnt pathway ligands, receptors, and antagonists (21,72–74). Pitx2 was found to interact with the promoters of five WNT ligands and induce expression of four of the ligands leading to activation of the β-catenin-dependent canonical Wnt pathway and enhanced cell proliferation in cancer cell lines (74). A zebrafish ortholog of one of the induced ligands, Wnt6, was also identified by our microarray; moreover, Wnt6 was also previously detected as a FOXC1 target in two independent assays (75) and thus may represent a common downstream target of two ARS factors. Also among the human orthologs of our microarray targets, WNT7A has been shown to induce limbal stem cells into skin-like epithelium when depleted (76), and an exonic variant in WNT10A has been implicated in decreasing central corneal thickness and doubling the risk of keratoconus in humans (77). While PITX2 was shown to exhibit a positive feedback control loop with WNT ligands, its overexpression was demonstrated to reduce the expression of several Frizzled receptors and lipoprotein receptor-related protein (LRP) co-receptors, thus suggesting that it also supplies negative feedback control to the pathway (74). Such negative regulation is consistent with the observation of the activation by Pitx2 of certain Wnt antagonists like Dkk2 (21). Further testing will be needed to verify if Pitx2 is a direct regulator of the identified Wnt ligands and antagonists or if the observed down-regulation of both in pitx2 mutants is a compensatory mechanism in which one class is directly targeted and the other fluctuates in response.

Another group of targets with significant down-regulation in pitx2 mutant eyes were transcripts encoding for collagens I and V. Collagen I is the primary component of the orthogonally arranged, stratified fibril layers that make up the vertebrate corneal stroma, and collagen V is a key support molecule for collagen I fibrill formation (43,78). Mutations in collagens have been identified as causative of Ehlers-Danlos syndromes (EDS), a genetically heterogeneous group of connective tissue disorders (47,79). In particular: mutations in COL1A1 (rare) and COL5A2 (major) are associated with classic EDS [MIM 130000] (79); mutations in COL1A1 and COL1A2 are associated with EDS, arthrodactylia type [MIM 130060] (79–81); and COL1A2 mutations are also associated with EDS cardiac valvular type [MIM 225320]. EDS exhibits several features that overlap findings present in ARS patients, including (but not limited to): microcornea; glaucoma; cornea plana; small, irregularly placed teeth; umbilical and inguinal hernias; mitral valve prolapse; and hypertensibility (47,82). Mutations in COL1A1 have been associated with various glaucoma phenotypes, including both primary open angle and infantile-onset/primary congenital; the majority of patients with these phenotypes also presented with osteogenesis imperfecta (OI), a group of inherited disorders characterized primarily by fragile bones and secondarily with mitral valve prolapse, blue sclerae, thin cornes, optic atrophy, vitreous hyperplasia and corneal opacities, among other features (49–51). Interestingly, one case of OI with features of both EDS and ARS has been attributed to a frameshift c.3313delA mutation in COL1A1 (52), further linking these two phenotypes. Finally, a partial deletion involving another collagen gene, COL5A3, and OLFM2 was recently identified in a patient with bilateral microphthalmia, sclerocornea, ptosis, small teeth, bifid uvula and hearing defects (46). Our work suggests that PITX2/pitx2 may regulate expression of genes encoding collagens I and V. With this, early deficiency in expression of corneal collagens is likely to contribute to the abnormal phenotypes associated with PITX2/pitx2 disruption. Further work will determine whether the down-regulation of collagen transcripts is a result of direct involvement of pitx2 in regulation of their expression or a secondary effect. Overall, the developed pitx2<sup>2Mel</sup> zebrafish line recapitulates the characteristic phenotypes of PITX2-mediated ARS. In addition to this, the possible role of PITX2 in corneal dermoid phenotypes is further supported by the presented zebrafish data. The initial molecular assessment of pitx2-deficient ocular tissues at early stages of development, prior to structural defects, identified significant changes in the expression of multiple Wnt and collagen genes, some of which have been previously associated with overlapping ocular and craniofacial phenotypes. The developed model will be useful in examining the mechanisms of a wide range of ocular phenotypes associated with PITX2 deficiency and identifying potential therapeutic targets.

**Materials and Methods**

**Human screening**

This human study was approved by the Institutional Review Board of Children’s Hospital of Wisconsin with written informed consent obtained for every subject including consent for photo publication. The diagnosis of ARS was based on clinical evaluation by referring physicians. In general, at least one family member needed to have characteristic ocular findings (posterior embryotoxon, iris processes, iris hypoplasia, pupillary anomalies) with two or more of the typical non-ocular features.
Electron microscopy was performed as described by Soules and Link (32). Briefly, 3- and 14-dpf embryos were collected and fixed in primary fixative (2% paraformaldehyde, 2.5% glutaraldehyde, 3% sucrose, 0.06% phosphate buffer (pH 7.4)) at 4°C. Specimens were submitted to the Electron Microscopy Facility at the Medical College of Wisconsin for embedding, which included 1-h post-fixation in 1% osmium tetroxide on ice for membrane preservation, dehydration through a series of methanol and acetone washes, and infiltration with EMB 812 resin (14120; Electron Microscopy Sciences, Hatfield, PA). Transverse, semi-thin (500 nm) sections were cut with a DiATOME Histo diamond knife (H10213; DiATOME, Hatfield, PA) on a PowerTome XL microtome (Boeckeler Instruments, Tucson, AZ) and collected throughout the central anterior chamber as defined by maximum lens diameter. Sections were heat-fixed to glass slides, stained with 1% Toluidine Blue in 1% Borax buffer, and imaged on an AxioImager.Z1 microscope with an AxioCam MRc5 camera and ZEN pro software (Zeiss, Thornwood, NY). Ultrathin sections (70–80 nm) were subsequently taken at ~3–5 μm intervals through the central anterior chamber with a DiATOME Ultra 45° diamond knife (MT7376; DiATOME), collected on copper hexagonal mesh coated grids (G200H-Cu; Electron Microscopy Sciences) and stained with uranyl acetate and lead citrate for contrast. Images were captured using a Hitachi H600 TEM microscope (Hitachi, Tokyo, Japan).

Inclusion body count and area measurements were performed with ImageJ software (https://imagej.nih.gov/ij/). Electron microscopy images of 3-dpf dorsal, central and ventral cornea for a wild-type and mutant embryo at the same magnification were used. Inclusion bodies were designated as electron-dense structures in the corneal surface epithelial and subepithelial layers that did not appear to be membrane-bound or within a membrane-bound organelle. All inclusion bodies identified in wild-type or mutant images in either surface or subepithelial layers were measured; the data from three images were pooled together and displayed as a box and whisker plot for each condition (wild-type versus mutant) for each epithelial layer. The statistical significance of average inclusion body area per layer between wild-type and mutant images was determined using the Student’s t-test.

For histology, 1-mpf and 1- to 2-year-old adult fish tissues were preserved in 10% formalin and submitted to the Children’s Research Institute Histology Core at the Medical College of Wisconsin for paraﬁn sectioning and H&E staining per standard protocols. Slides were imaged on a NanoZoomer digital slide scanner, and images were viewed with NDP.view2 viewing software (Hamamatsu, Hamamatsu City, Japan).

PAS staining was performed using standard protocols. 1-mpf and adult paraﬁn sections were immersed in 1% acetic acid [hydrochloric acid and 70% alcohol] for 5 min to de-colorize following H&E staining. Slides were oxidized in 0.5% periodic acid for 5 min, rinsed in distilled water, and stained with Schiff’s reagent (Newcomer Supply, Middleton, WI) for 15 min. Sections were rinsed in warm tap water for 7 min then in distilled water before being counterstained with hematoxylin for 1 min and rinsed again. Slides were placed in clarifier for 30 s then rinsed 1 min each in distilled water, 0.1% ammonium water, and distilled water; they were quickly dehydrated through two changes of 95% alcohol, three of 100% alcohol and three of xylene before cover-slipping and imaging as described earlier.

For immunohistochemistry, 1-mpf and 1- to 2-year-old adult paraﬁn sections on cover-slipped slides were soaked in xylene for 2–3 days to remove the coverslip. Paraﬁn sections

Characterization of embryonic and adult phenotypes in pitx2 mutants

Alcian Blue cartilage staining was performed as per Barral-Gimen et al. (84). Briefly, embryos were collected at 5-dpf and fixed overnight in 4% paraformaldehyde in 1 x diethyl pyrocarbonate (DEPC)-phosphate-buffered saline (PBS) at 4°C. Following two PBS washes, embryos were bleached in 1 ml 10% hydrogen peroxide (H2O2) supplemented with 50 μl of 2M KOH for 1 h on a rotating platform. Embryos were stained overnight in 0.1% Alcian Blue solution comprised of Alcian Blue 8GX (Sigma-Aldrich, St Louis, MO) dissolved in acidic ethanol. Stained embryos were washed thoroughly with acidic ethanol, digested in protease K for 1 h, re-fixed in 4% paraformaldehyde, progressively dehydrated with ethanol, and stored in 80% glycerol at 4°C. Craniofacial structures were identiﬁed as presented by Piotrowski et al. (85). Whole mount images were obtained on a Nikon SMZ1500 dissection microscope (Nikon Instruments Inc., Melville, NY).
were exposed to antigen retrieval for 10–20 min at 100 °C then kept at room temperature for 20-min followed by two 5-min water rinses and one 10-min PBS plus 0.1% TWEEN-20 (PBT) wash. Slides were blocked in 10% donkey serum for 2 h and incubated in primary antibody overnight at 4 °C. Slides underwent two 10-min PBT washes followed by a 2-h incubation in secondary antibody and imaging as previously described. Primary antibodies utilized were collagen Iα1 (col1α1a; 1:100; GTX124368; GenTex, Inc., Irvine, CA), CKS proteoglycan (1:200; MAB2022; EMD Millipore, Billerica, MA), N-cadherin (cdh2; 1: 50, GTX125962; GenTex) and DAPI (1:1000; 62247; Thermo Fisher Scientific, Waltham, MA). Secondary antibodies used were donkey anti-rabbit Alexa Fluor 488 IgG (1:100; A-21206; Thermo Fisher) and donkey anti-mouse Alexa Fluor 568 IgG (1:100; A-10037; Thermo Fisher).

For ISH and TUNEL staining, embryos underwent standard fixation protocols. Embryos were collected at the desired stage, washed twice with 1× DEPC-PBS and fixed overnight in 4% paraformaldehyde in DEPC-PBS at 4 °C. Embryos underwent two 5-min ice cold PBS washes and four 5-min ice cold 100% methanol washes then were stored at −20°C.

ISH was performed as previously described in (37). A foxc1a plasmid was ordered from Open Biosystems (8001422; GE Healthcare, Piscataway, NJ) and linearized with EcoRI (New England Biolabs). An 843-bp foxc1a digoxigenin-labeled antisense RNA probe was generated using the DIG RNA labeling mix from Roche Diagnostics and 177 RNA polymerase (Roche Diagnostics Corporation, Indianapolis, IN). Whole mount embryos were imaged as described above.

RNAscope probe Dr-Pitx2 (467531-C1; Advanced Cell Diagnostics, Newark, CA) was obtained and used as described by Gross-Thebing et al. (86). For cryosectioning, stained embryos were prepared by undergoing: two PBS washes; two overnight infiltrations at 4 °C; first with 30% sucrose/PBS then with TissueTek O.C.T. Compound (Sakura Finetek USA, Inc., Torrance, CA); embedding in cryomolds; and freezing. 10-micron sections were melted onto charged glass slides, mounted with ProLong Gold embedding in cryomolds; and freezing. 10-micron sections were stained as described above.

For TUNEL staining, embryos were fixed in 4% paraformaldehyde for 10–15 min, washed twice with PBS, and stained using a TUNEL reaction kit (Roche Diagnostics). Embryos were imaged as described above.

**Transcriptome analysis in pitx2M644+ embryos**

To collect embryonic ocular tissues from mutant and wild-type embryos, whole eyes of 23-hpf embryos were dissected while the remaining tissue was utilized for genotyping. Wild-type and pitx2M644 homozygous mutant whole eyes were then pooled together for RNA extractions (final samples contained 56, 58 and 50 eyes for homozygous mutants and 88, 109 and 112 for wild-type tissues). RNA was extracted from the collected eyes using TRIzol and the Direct-zol RNA MiniPrep Kit (Zymo Research, Irvine, CA). RNA quality was assessed on both a NanoDrop 2000 UV-Vis Spectrophotometer (Thermo Fisher) and an Agilent 2100 Bioanalyzer utilizing the Agilent RNA 6000 Pico Kit (Agilent Technologies, Santa Clara, CA). RNA from three independent pitx2M644 homozygous eye collections with RNA Integrity Numbers (RINs) of 8.9, 8.6 and 9.0 and three independent wild-type collections with RINs 8.7, 8.9, and 8.7 was submitted to OakLabs (Hennigsdorf, Germany) for transcriptome analysis using an ArrayXS Zebrafish v1 microarray containing 60 023 target IDs. Statistical analysis of the microarray data was performed by Oaklabs and consisted of quantile normalization using ranked median quantiles, conversion to log2 values for calculation of control versus mutant means and standard deviations per target, and Welch’s t-test (or unequal variances t-test) that was used to test the hypothesis that the signals of the gene in the two biological groups of samples (e.g. mutant and wild-type) have equal means. Differentially expressed targets were defined as having a change in mean log2 >1 between mutant and wild-type values, correlating to a 2-fold change in expression, and a P-value < 0.05. For pathway analysis, zebrafish-specific transcripts were annotated with human orthologs using Ensembl Biomart (http://useast.ensembl.org/biomart). Remaining transcripts with no identified orthologs were manually annotated by using the provided Ensembl transcript id to search for orthologs or paralogs within Ensembl [http://grch37.ensembl.org/Danio_rerio (Zebrafish Zv9); http://useast.ensembl.org/Danio_rerio (Zebrafish GRc10) or to perform a protein BLAST (https://blast.ncbi.nlm.nih.gov). The list of human orthologs was analyzed for pathway enrichment by Ingenuity Pathway Analysis software (Qiagen, Hilden, Germany). Microarray data were submitted to the ArrayExpress database at EMBL-EBI under Accession Number E-MTAB-6463 (www.ebi.ac.uk/arrayexpress).

Select identified targets were verified by real-time quantitative PCR using transcript-specific primers (Supplementary Material, Table S1), SYBR Green PCR Master Mix (Applied Biosystems, Waltham, MA) and CFX Connect and CFX96 Touch Real-Time PCR Detection Systems (Bio Rad, Hercules, CA). 24-, 48- and 72-hpf wild-type and pitx2M644 homozygous eye cDNA was generated from equal concentrations of wild-type and pitx2M644 homozygous eye cDNA utilizing SuperScript III reverse transcriptase (Thermo Fisher). Each experiment included the following conditions at 200 pg cDNA per technical replicate: 24-, 48- and 72-hpf wild-type eye cDNA; 24-, 48- or 72-hpf pitx2M644 homozygous eye cDNA; no reverse transcriptase control; and no template control which excluded primers. All samples were
normalized to β-actin and run in triplicate to obtain average Cq values. Total fold changes and standard deviations were calculated as the average of three independent biological repeats via the efficiency-corrected ΔCq model, which accounts for differing primer efficiencies. Statistical significance was determined using the Student’s t-test.

For heatmap generation, the collective average and standard deviation of the normalized three wild-type and three pitx2<sup>1692</sup>-deletions replicates for each target of interest were calculated, and the Z-score for each replicate was derived by dividing the difference between the replicate and the mean by the standard deviation. Z-scores were imported into the SNP and Variation Suite software (Golden Helix, Inc., Bozeman, MT) for heatmap synthesis.

**Supplementary Material**

Supplementary Material is available at HMG online.

**Acknowledgements**

The authors gratefully acknowledge the patients and their families for their participation in research studies. We would also like to thank the Department of Cell Biology, Neurobiology and Anatomy for use of shared equipment, as well as Christine Duris and the Children’s Research Institute Histology Core and Clive Wells and Robert Goodwin from the Medical College of Wisconsin Electron Microscopy Core Facility for technical assistance.

Conflict of Interest statement. None declared.

**Funding**

This work was supported by awards from the National Institutes of Health (EY015518) as well as funds provided by the Children’s Research Institute Foundation at Children’s Hospital of Wisconsin (E.V.S.), NEI Training Grant (5T32EY014537-13 and -14), and NIH Core Grant (P30EY001931). Funding to pay the Open Access publication charges for this article was provided by the Department of Pediatrics at the Medical College of Wisconsin.

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