Hypo- and Hypermorphic FOXC1 Mutations in Dominant Glaucoma: Transactivation and Phenotypic Variability

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Abstract

Dominant glaucoma, a heterogeneous, infrequent and irreversible optic neuropathy, is often associated with elevated intraocular pressure and early-onset. The role of FOXC1 in this type of glaucoma was investigated in twelve Spanish probands via nucleotide variation screening of its proximal promoter and unique exon. Functional evaluations of the identified variants included analyses of the transcriptional activity, protein stability, DNA binding ability and subcellular localization. Four different mutations that were identified in four probands (33.3%) were associated with remarkable phenotypic variability and were functionally classified as either hypermorphic (p.Y47X, p.Q106X and p.G447_G448insDG) or hypomorphic (p.I126S) alleles. To the best of our knowledge, three of the variants are novel (p.Y47X, p.I126S and p.G447_G448insDG) and, in addition, hypermorphic FOXC1 mutations are reported herein for the first time. The presence of an intact N-terminal activation domain in the truncated proteins p.Y47X and p.Q106X may underlie their associated transactivation hyperactivity by a gain-of-function mechanism involving dysregulated protein-protein interactions. Similarly, altered molecular interactions may also lead to increased p.G447_G448insDG activity. In contrast, the partial loss-of-function associated with p.I126S was due to impaired protein stability, DNA binding, protein phosphorylation and subcellular distribution. These results support that moderate and variable FOXC1 transactivation changes are associated with moderate goniodysgenesis, dominant glaucoma and remarkable phenotypic variability.
Introduction

Glaucoma is a highly heterogeneous, irreversible and progressive blinding disease produced by the death of the retinal ganglion cells, which results in the degeneration of the optic nerve. Raised intraocular pressure (IOP) is the primary risk factor for developing glaucoma. Most glaucoma cases behave as a late-onset complex disease and a minority of cases show an early-onset and follow simple Mendelian inheritance. Dominant transmission is typically observed in juvenile open-angle glaucoma (JOAG; MIM# 137750), as well as in some cases of both adult-onset primary open-angle glaucoma (POAG; MIM# 137760) and primary congenital glaucoma (PCG; MIM# 231300) [1, 2]. PCG is caused by developmental abnormalities in the anterior segment of the eye, which is required for aqueous humour drainage, and manifests clinically in the neonatal or infantile period, generally before the age of 3. Moreover, late congenital glaucoma cases (LCG, i.e., glaucoma diagnosed over 3 years of age with abnormal gonioscopy of the anterior segment and dominant inheritance) have also been described [3]. So far two primary genes, CYP1B1 [4] (MIM# 601771) and LTBP2 [5] (MIM# 602091), have been identified in recessive PCG. Loss-of-function CYP1B1 mutations are the predominant known genetic cause of this type of glaucoma in different populations [6–8]. MYOCILIN (MYOC, MIM# 601652) [9, 10] and FORKHEAD BOX C1 (FOXC1 MIM# 601090) alterations have also been found in a small number of PCG patients [3, 10]. JOAG presents with an early age of onset, usually between 10 and 35 years. Although at least five loci have been identified for this type of dominant glaucoma, only one disease-causing gene (MYOC) has been identified thus far [11]. MYOC mutations are present in approximately 10% of all JOAG cases [12]. Finally, nine dominant adult-onset POAG loci have been reported, but only three genes related to this type of glaucoma have been identified, OPTN (MIM# 602432) [13], ASB10 (MIM# 615054) [14] and WDR36 (MIM# 609669) [15].

FOXC1 lies in the 6p25 forkhead cluster (FOXC1/FOXF2/FOXQ1) and mutations in this gene cause a spectrum of autosomal dominant anterior eye segment defects, including Axenfeld-Rieger syndrome type 3 (ARS; MIM# 602482), causing an increased risk for glaucoma and varying degrees of iris or extra-ocular abnormalities [16]. This gene encodes a member of the FOX class of transcription factors, which are involved, among other processes, in the regulation of craniofacial, cardiovascular and ocular development [17]. A characteristic and conserved 110-amino acid DNA-binding domain, known as the forkhead domain (FHD), is present in the protein. FOXC1 is expressed in mesoderm and neural-crest-derived cells, including the cells in the anterior segment of the eye, and the periorcular mesenchyme and mesenchymal cells that have migrated into the eye [18, 19]. The transcriptional activity of this phosphoprotein is regulated by N- and C-terminal activation domains [20]. To date, more than fifty different missense, nonsense, and frameshift FOXC1 mutations have been identified, with the majority affecting the forkhead domain [21]. These mutations reduce the FOXC1 transactivation ability [22], and none have been described to increase FOXC1 activity. However, increased activity is likely to be associated with duplications in the gene, which has been observed in various types of anterior segment disorders and glaucoma [23]. It has been shown that patients diagnosed with Axenfeld-Rieger malformation who carry FOXC1 duplications have a more severe prognosis for glaucoma development than patients with FOXC1 mutations [24].

To our knowledge, this is the first study to identify hypermorphic FOXC1 mutations, and to show that moderate and variable residual FOXC1 activity levels are involved in dominant glaucoma, and may contribute to the phenotypic variability that is present in this disease.
Materials and Methods

Ethics statement

The study and informed consent procedures were approved by the Ethics Committee for Human Research of the Hospital Clínico San Carlos, Madrid (Spain), and followed the tenets of the Declaration of Helsinki. Informed written consent was obtained from all of the participants and was recorded by staff involved in the study.

Subjects

Twelve unrelated probands affected by dominant glaucoma were included in this study. All subjects were clinically evaluated by glaucoma specialists. The ophthalmic examination included slit lamp biomicroscopy, gonioscopy, biometry, intraocular pressure (IOP) measurement and ophthalmoscopy. The PCG clinical diagnosis included at least two of the following clinical features; increased corneal diameter (>12 mm), along with elevated IOP (>21 mmHg or >16 mmHg under general anaesthesia with sevoflurane) and/or Haab’s striae, corneal edema and optic disc changes, usually before the age of 3. The preoperative PCG diagnosis was based on early clinical signs and symptoms (photophobia, blepharospasm, tearing, corneal edema and corneal enlargement). Patients over 3 years of age who were diagnosed with abnormal gonioscopy of the anterior segment (high iris insertion and absence of an angle recess), were considered to be LCG cases. Cases diagnosed between 10 and 35 years of age or over the age of 40, were classified as JOAG and POAG, respectively. All of the following criteria were required for either the JOAG or POAG diagnosis: open anterior chamber angle, IOP > 21 mm Hg, characteristic optic disc changes (e.g., vertical cup-to-disc ratio > 0.4, thin or notched neuroretinal rim or disc haemorrhage), and characteristic visual field changes. The visual field alterations were classified as previously described [25]. Secondary glaucoma probands were excluded from the study. In children, the intraocular pressure was measured using the Perkins applanation tonometer (Clement Clarke MK2, Harlow, UK), generally under sevoflurane anaesthesia; and as soon as they were sufficiently anesthetized to check the intraocular pressure (during the first 10 min after induction and measured again immediately before the children wake up when they breathe air). Older and more cooperative children were assessed using a slit lamp for tonometry with a Goldmann applanation tonometer.

Mutation screening

Genomic DNA was extracted from the peripheral blood, using the QIAamp DNA Blood Mini Kit (Qiagen). The DNA sequence variation analyses were carried out using automatic Sanger sequencing. The promoter (nucleotides −1 to −875), the translated region and the 5’- and 3’-untranslated regions (UTRs) of FOXC1 were amplified via PCR using the primers described in S1 Fig. and S1 Table. The mutations were confirmed in two independent PCR amplifications by DNA sequencing.

Site directed mutagenesis and cloning of the identified FOXC1 mutations

Three FOXC1 mutations (p.Y47X, p.Q106X and p.I126S) and the control FOXC1 mutation p. I126M were obtained via site-directed mutagenesis using the QuickChange Site-directed Mutagenesis Kit (Stratagene), with the primers and PCR conditions indicated in the S1 Materials and Methods and S2 Table. The p.G447_G448insDG variant was directly amplified from the genomic DNA of the carriers using primers P1 and P2 (S1 Materials and Methods), and were cloned into the EcoRI/BamHI restriction sites of the pcDNA3.1(-) vector. All of the recombinant FOXC1 versions were transiently expressed in human embryonic kidney 293T
HEK-293T cells purchased from the American Type Culture Collection (ATCC Rockville, MD) as previously described [26, 27] and indicated in the S1 Materials and Methods. Transient plasmid transfections were carried out using 50–500 ng of plasmid DNA using the Superfect Transfection Reagent (Qiagen), according to the manufacturer’s instructions.

Transcriptional activity assays

The FOXC1 transactivation assays were performed using the Luciferase Assay System (Promega) according to the manufacturer’s instructions. A 600-bp fragment of the human CXCR4 gene promoter, which contains one FOX-binding element (FBE) [28], was cloned into the Nhel/NcoI restriction sites of the pGL3-basic vector (Promega) via directional PCR using normal human genomic DNA as a template and the following primers: 5'TCTGGCTAGCCGCGGGGAATGGCGTTGG3 and 5'CTCCATGGTAACCGCTGGTTCTCCAG3 (the Nhel and NcoI sequences are indicated with bold letters, respectively). The monkey kidney derived COS7 cell line has been widely used in functional analyses of FOXC1 mutations associated with human ocular phenotypes [29, 30]. In this study, we employed the human embryonic kidney 293T cell line (HEK-293T). HEK-293T cells in 24-well tissue culture plates (2.5x10⁵ cells/well) were transfected with 500 ng of the FOXC1 expression vector, along with 50 ng of the recombinant pGL3-basic-CXCR4 luciferase reporter, and 50 ng of the pMirTarget vector (Origene), which expresses red fluorescent protein (RFP) as a transfection control. The total amount of transfected DNA was equalized to the empty vector. The transactivation assays were performed 48 h after the transfection. The cells were harvested and assayed for firefly luciferase activity as previously indicated. FOXC1, RFP and LDH were detected by Western blot as the expression, transfection and loading controls, respectively. The RFP and LDH bands were quantitated via densitometry (n≥3) and significant differences were analyzed using the t-test.

Protein stability and half-life calculation

The protein stability was studied via western blot of the transfected cells incubated with cycloheximide (300 μg/ml) at different times, as previously reported [31]. The FOXC1 protein levels were determined by densitometry and the relative amounts at the different times after cycloheximide treatment were expressed as a percentage of the levels at time 0 h. At least three independent assays for each variant were performed. The transfection efficiency was assessed by co-transfecting the cDNAs encoding the different mutants (500 ng) with the pMirTarget vector (50 ng), which encodes RFP. The fluorescent protein was detected via western blotting. The loaded samples were normalized for total protein content using Bradford reagent (Pierce). FOXC1 decay follows a first-order kinetics. The slope of the decay line was calculated using standard linear regression, and the protein half-life was determined as previously described [32].

Nuclear protein extraction and Electrophoretic Mobility–Shift Assay (EMSA)

The nuclear extracts of the HEK-293T cells expressing recombinant FOXC1 proteins were prepared as previously reported [22] and as briefly described in the S1 Materials and Methods. An EMSA was performed using the LightShift EMSA Optimization and Control Kit (Thermo Scientific) according to the manufacturer’s protocol and as described in the S1 Materials and Methods.
FOXC1 treatment with calf intestinal alkaline phosphatase

The phosphorylation status of the FOXC1 protein expressed in the HEK-293T cells was analyzed by calf intestinal alkaline phosphatase (CIP) treatment of the nuclear extracts. The nuclear extracts were incubated with either 5 units of CIP, 11 \( \mu \text{m} \) sodium vanadate (NaVO\( _3 \)), or both for 2 h at 37°C. The recombinant proteins were detected via western immunoblot using an anti-myc antibody. The percentage of phosphorylated protein was estimated via densitometry of western blot bands as follows: \[ \text{percentage of phosphorylated protein} = \frac{\text{upper band volume (phosphorylated FOXC1)} \times 100}{\text{upper band volume (phosphorylated FOXC1)} + \text{lower band volume (dephosphorylated FOXC1)}} \]. LDH was detected via western blot as the loading control, and was quantitated via densitometry \( (n \geq 3) \), and significant differences were analyzed using the \( t \)-test.

Western and dot blotting and antibodies

For the western blot analysis, the nuclear extracts were fractionated using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) using the Mini-PROTEAN III Gel Electrophoresis System (BioRad). A tricine-SDS-PAGE was performed as previously described \[33\]. The protein content of the samples were normalized using the Bradford assay. The gels were subsequently transferred onto Hybond ECL nitrocellulose membranes (Amersham) for the immunodetection, And a dot blot analysis of the non-sense mutants was carried out by spotting the normalized nuclear extracts onto Hybond ECL nitrocellulose membranes in a blot transfer apparatus (Bio-Dot, Bio-Rad). A commercial mouse monoclonal anti-myc (Santa Cruz Biotechnology) antibody was used as the primary antibody, diluted at 1:1000. Horse-radish peroxidase-conjugated antibodies against mouse IgG (Pierce) were diluted to 1:1000–1:4,000. The chemiluminescence detection was performed using Supersignal Dura Western Blot reagents (Pierce), and luminescent imaging was used for the chemiluminescence detection (LAS3000-mini, Fujifilm, Tokyo, Japan). The densitometry for the protein band quantitation (FOXC1, LDH and RFP) was performed using the Quantity One 4.1 analysis software package (BioRad) in at least two independent experiments performed in triplicate. Ponceau S (Panreac) staining of the blots prior to the antibody incubation was performed to ensure the integrity of the samples and that equal amounts of the sample were analyzed \[34\]. As an additional sample loading control, lactate dehydrogenase (LDH) was detected in the cell extracts and culture media using a goat anti-LDH antibody [AB1222, Chemicon, diluted to 1:5000] and an anti-goat IgG horse-radish peroxidase-conjugated antibody (sc-2033, Santa Cruz Biotechnology, diluted to 1:2000). The RFP (transfection control) was detected using a rabbit anti-RFP antibody (#AB233, Evrogen), diluted to 1:5000, and an anti-rabbit IgG horse-radish peroxidase-conjugated antibody (#1858415, Pierce) diluted to 1:1000.

Immunocytochemistry

The HEK-293T cells were seeded on coverslips placed into 24-well plates, and were transiently transfected with DNA constructs encoding different FOXC1 variants. All of the transfections were performed using the SuperFect Transfection Reagent (Qiagen). After the transfection, the cells were washed once with DPBS and cultured for 24 h. The cells were then fixed with 4% paraformaldehyde for 10 min at room temperature, followed by incubation with phosphate-buffered solution containing 0.2% triton X-100, 10% FBS, and 5% bovine serum albumin for 30 min at room temperature. The recombinant proteins were detected using an anti-myc antibody (Santa Cruz Biotechnology) (at a 1:500 dilution) at 4°C overnight followed by a Cy3-conjugated anti-mouse IgG (Jackson ImmunoResearch Labs,1:1000 dilution) for 2 h at room temperature. Finally, the coverslips were mounted on glass slides using polyvinyl alcohol
mounting medium with DABCO (Fluka) containing DAPI (4',6'-diamidino-2-phenylindole; F6057 SIGMA) and were viewed under a laser scanning confocal microscope (Zeiss LSM 710).

**Bioinformatic analyses**

The *In silico* analyses of the sequences of the different *FOXC1* variants were carried out using the programs described in the S1 Materials and Methods. Mutations were named according to RefSeq: NM_001453.2 and using the directions from Mutalyzer (https://humgenprojects.lumc.nl/trac/mutalyzer). The transcription start site and 5'-UTR sequence were defined as previously described [35]. The first nucleotide of the translation initiation site was numbered as nucleotide +1. The novel variants have been submitted to dbSNP (http://www.ncbi.nlm.nih.gov/SNP/).

**Statistical analyses**

The statistical comparisons between the groups were performed using either the *t*-test or the one-way analysis of variance (ANOVA). A Bonferroni correction was applied to adjust the tests for multiple comparisons. The data were statistically processed using the SigmaStat 2.0 software (SPSS Science).

**Results**

**The identification of rare *FOXC1* mutations**

To analyze the role of *FOXC1* mutations in dominant glaucoma, a total of twelve unrelated probands with no mutations in the glaucoma genes MYOC and CYP1B1 were studied. Four of them were affected with PCG, one with LCG, three with JOAG and four with POAG. Four different rare heterozygous sequence variations were identified in four probands (33.3%; two PCGs, one LCG and one POAG; Fig. 1 and Table 1). Three of these variants (p.Y47X, p.I126S and p.G447_G448insDG) were not found in the Emsembl variation database, the Human Gene Mutation Database, GeneCards, OMIM or in pubMed, and were therefore considered to be novel mutations. These mutations were also absent in 200 control chromosomes (data not shown). The mutations p.Y47X and p.I126S segregated with the disease in families PCG68 and PCG73, respectively (Fig. 1A and B). These two families showed remarkable phenotypic variability, particularly, but not exclusively, for the disease onset. Proband PCG68, who was classified as LCG, showed a thin iris stroma and posterior embrytoxon in both eyes and no evident iris hypoplasia, corectopia or systemic alterations (Fig. 2A and B). Her mother, diagnosed with JOAG at the age of 30, also exhibited posterior embryotoxon with no iris attachments or areas of iris atrophy secondary to surgery (Fig. 2C). Generalized stromal thinning, atrophy, corectopia (with the pupil pulled in the direction of the most prominent angle findings), or iris hole formation occurring in the quadrant opposite the site of corectopia, were not observed in these two patients; therefore, according to the 9th Consensus Report of the World Glaucoma Association, the presence of mild iris stroma anomalies in the proband were considered to be compatible with congenital glaucoma [36]. Proband PCG73 was diagnosed with congenital glaucoma and her mother was diagnosed with juvenile glaucoma. Both patients presented thinned iris stromal areas, posterior embryotoxon with no iris attachments, surgical iridotomies in both eyes and normal pupils (Fig. 2D-G). Generalized iris stromal thinning, atrophy, corectopia or iris hole formation were also not detected in these patients. Interestingly, another PCG73 family member, subject II:4, also carried the same *FOXC1* genotype and presented corectopia, generalized iris stromal thinning, which could be secondary to surgery, posterior embryotoxon and surgical iridectomy in his left eye (Fig. 2H). He did not show dental, umbilical (Fig. 2I and J, respectively) or any other systemic malformations, and was diagnosed with Axenfeld-Rieger...
Fig 1. FOXC1 mutation segregation in families with autosomal dominant glaucoma. (A, B, C and D) Pedigrees of the families. The numbers in the inferior left side of the symbols indicate age at diagnosis. The oblique lines represent dead subjects. Black, dark grey and light grey symbols indicate glaucoma, Axenfeld-Rieger anomaly and ocular hypertension, respectively. The arrows in the pedigrees show the index cases. +: wild-type allele. The insets in each panel correspond to the electropherograms of the FOXC1 mutations identified in each family. To facilitate the comparison, the wild-type sequence is...
anomaly (ARA). His right eye was eviscerated at the time of the study. In family 559–10, the proband was the only subject available for genetic analysis, and ocular photographs were not available. She was diagnosed with PCG and carried the nonsense mutation p.Q106X, (Fig. 1C and Table 1). Finally, in the adult-onset glaucoma family (556–10), the proband was diagnosed at the age of 66, and her three siblings manifested glaucoma in their fifties (Fig. 1D and Table 1). Only the index case and her sister consented to the genetic analyses and were found to carry the heterozygous FOXC1 variant p.G447_G448insDG (Fig. 1D, subjects II:2 and II:3). The proband’s iris and iridocorneal angle were normal (Fig. 2K-M) and a careful clinical examination directed to detected Axenfeld-Rieger alterations did not reveal any other relevant ocular or systemic anomalies. All of the described mutations are associated with bilateral glaucoma, variable iris morphology and a disease onset ranging from 1 to 66 years in the index cases (Table 1). The previously described 3’-UTR variant c.734A>T (rs35717904) was also identified in probands PCG73 and 556–10 (data not shown). PITX2 mutations were ruled out in these patients.

To evaluate the pathogenicity of the identified variants a number of approaches were employed, including bioinformatic analysis, evolutionary conservation of the mutant amino acids or nucleotides and an in vitro assessment of transcriptional activity, protein stability, DNA binding ability, phosphorylation status and subcellular localization.

Bioinformatic analysis and evolutionary conservation of the variants

The analyses of the variants with the programs indicated in the S1 Materials and Methods showed that three coding mutations mapped to activating domain 1 (p.Y47X), alpha helix-2 (p.Q106X) and alpha-helix 3 (p.I126S) of the forkhead DNA binding domain (Fig. 3A). The shown above the corresponding mutant sequence. All the mutations were detected in the heterozygous state. Arrows in the electropherograms indicate the location of mutations. Duplicated nucleotides in panel D inset are indicated by boxes.

| Proband or family member | Nucleotide change | Amino acid change | Phenotype/lat | Age at diagnosis (years/sex) | IOP (mmHg) at diagnosis (OD/OS) | C/D (OD/OI) | Treatment |
|--------------------------|-------------------|------------------|--------------|----------------------------|---------------------------------|------------|-----------|
| PCG68 (III:1)            | c.141C>G          | p.Y47X           | LCG/B        | 7/F (24/20)                 | NA                              |            | 1         |
| PCG68 (II:4)             | c.141C>G          | p.Y47X           | JOAG/B       | 30/F 30/25                 | 0.8/0.8                        | Surgery    | 5         |
| 559–10 (II:1)            | c.316C>T          | p.Q106X          | PCG/B        | 3/F (25/25.9)              | NA                              |            | NA        |
| PCG73 (III:1)            | c.377T>G          | p.I126S          | PCG/B        | 1/F (14/15)                | (0.6–0.7/0.8–0.9)              | 1+Surgery  | 1+Surgery |
| PCG73 (II:2)             | c.377T>G          | p.I126S          | JOAG/NA      | 12/F                       | NA                              |            | NA        |
| PCG73 (II:4)             | c.377T>G          | p.I126S          | ARA/B        | NA/M                       | NA                              |            | NA        |
| 556–10 (II:2)            | c.1337_1342dup    | p.G447_G448insDG| AOG/B        | 66/F (21/20)               | NA                              |            | 1         |
| 556–10 (II:3)            | c.1337_1342dup    | p.G447_G448insDG| AOG/B        | >50/F                      | NA                              |            | NA        |

1The variants were present in the heterozygous state; 2Number of drugs and/or surgery; 3Novel variants; AOG/ARA/JOAG/LCG/PCG: Adult-onset glaucoma/Axenfeld-Rieger anomaly/juvenile-onset glaucoma/late-onset primary congenital glaucoma/primary congenital glaucoma; B: bilateral; C/D: cup/disk ratio; NA: Not available; OD/OS: right eye/left eye. Mutations were named according to RefSeq: NM_001453.2 and using directions from Mutalyzer (https://humgenprojects.lumc.nl/trac/mutalyzer).
remaining coding mutation (p.G447_G448insDG) was situated in a Gly-rich sequence that is part of the inferred intrinsically disordered region 4 (IDR4, Fig. 3A). An in silico analysis carried out using the SIFT, PolyPhen-2 and Panther programs to evaluate the functional effect of the amino acid substitution p.I126S predicted that it was damaging (SIFT score = 0.01;
PolyPhen-2 score = 0.99; Panther Pdeleterious = 0.99). The evolutionary conservation analysis revealed that the amino acid residue I126 is conserved from fish to mammals and that the DG insertion affects a poly-Gly tract that is shared only by humans and chimpanzees (Fig. 3B).

Transcriptional activity

The transcriptional activity of the recombinant FOXC1 variants was assessed via a transient co-transfection with the luciferase reporter gene coupled to the CXCR4 promoter region, which contains one FBE (Fig. 4A). The detection of the truncated proteins p.Y47X and p.Q106X via standard SDS-PAGE and western blotting was prevented due to their small molecular size. To overcome this difficulty, the electrophoresis was carried out in the presence of tricine. Under these conditions, only p.Q106X (but not p.Y47X) could be identified (S2 Fig.). We also observed that p.Q106X and the wild-type protein were transferred into the nitrocellulose membrane with different yields due to their different molecular weights, making the quantification challenging. Because of these limitations, the truncated proteins were finally analyzed via dot-blot. Western or dot-blot analyses of RFP and LDH were used as controls for the transfection efficiency and sample loading, respectively, and revealed no significant differences. Therefore, the observed FOXC1 band intensity variations can be attributed to alterations in the protein stability, which will be shown later. Three mutations (p.Y47X, p.Q106X and p.G447_G448insDG) showed significantly increased transcriptional activity, ranging from approximately 160% to almost 200% of the wild-type protein, and were classified as hypermorphic variants (Fig. 4B and C). In contrast, the activity of the fourth variant (p.I126S) was only 20% of the normal protein and was similar to that of the control mutation p.I126M (Fig. 4C), showing that it is a hypomorphic allele. p.I126M has been identified in patients with ARS and glaucoma [16] and it has been proposed to be a positive control mutant for activity studies in the forkhead family of transcription factors [22].
Fig 4. Altered transcriptional activity of the FOXC1 variants identified in dominant glaucoma patients. (A) Scheme of the cDNA construct containing a FOXC1-binding element present in the CXCR4 promoter fused to the luciferase coding region. The cDNA was cloned into the PGL3 basic vector. This construct was used as a reporter of transcriptional activity in co-transfection assays with the different FOXC1 variants and the empty pMirTarget vector, which encodes RFP (transfection efficiency control). The numbers below the scheme correspond to nucleotide positions. cDNA constructs encoding the indicated FOXC1 variants (B) were transiently co-expressed with the reporter cDNA in HEK-293T cells. The transcriptional activity, expressed as a percentage of the luciferase activity of the wild-type protein, was measured as indicated in the Materials and Methods. The protein levels of the different FOXC1 versions present in HEK-293T cells 48 h after transfection were determined by western immunoblot using a monoclonal anti-myc antibody (Santa Cruz), except for the two non-sense mutants (p.Y47X and p.Q106X) which were difficult to detect and quantify by this technique and had to be assessed by dot-blot. Each lane contained 15 μg of total protein obtained from the cell lysates. Transfection efficiency was assessed via western immunoblot using an anti-RFP antibody (Evrogen). The sample loading control, endogenous LDH, was also detected via immunoblot using an anti-LDH antibody (Chemicon). Error bars correspond to the SD of three independent experiments carried out in triplicate. insDG: p.G447_G448insDG. NT: non-transfected cells (negative control). Asterisks indicate statistical significance as compared to the control: p < 0.05 (*); p < 0.01 (**); p < 0.001 (**). Significance was calculated by one-way ANOVA followed by Tukey multiple-comparison test.
Protein stability

The protein stability of the FOXC1 mutants was assessed in transiently transfected cells treated with the protein synthesis inhibitor cycloheximide. The FOXC1 levels were evaluated using either dot blot or western blot at different times after the inhibition of the protein synthesis. Any significant transfection or loading sample variations were ruled out via a western blot analysis of RFP and LDH, respectively (Fig. 5A). p.I126S showed remarkably reduced stability (Fig. 5A and B), with an estimated half-life value that was approximately 3-fold lower than that of normal FOXC1 (4.35 h vs. 12.95 h, respectively, Fig. 5C), and almost identical to the half-life of the control mutation p.I126M (4.20 h, Fig. 5C). The dot-blot stability analyses of the two dominant nonsense mutations (p.Y47X and p.Q106X) showed the shortest half-lives among all of the mutants (< 2.4 h).

p.I126S phosphorylation

A careful western immunoblot analysis of FOXC1 revealed a doublet with a low molecular weight band that was more intense in p.I126S than in the wild-type protein (Fig. 6A, lanes 5 and 1, respectively). To assess whether this difference is due to phosphorylation, p.I126S was
treated with CIP and analyzed via western blot. As controls, wild-type FOXC1 and the p.I126M mutant were treated in parallel. The LDH detection via western blot showed no significant sample loading variations (Fig. 6A). The untreated wild-type FOXC1 presented an upper band, estimated to represent 80% of the total protein (Fig. 6A, lane 1 and Fig. 6B). The proportion of this band in the untreated p.I126S (Fig. 6A, lane 5) represented less than 60% of the mutant protein (Fig. 6B), whereas in the control mutant was similar to the wild-type (Fig. 6A, lane 9 and Fig. 6B). The CIP treatment of all of the variants increased the lower band proportion (Fig. 6A, lanes 2, 6 and 10), and this effect was counteracted with the CIP inhibitor NaVO₃.

**Fig 6.** The p.I126S mutation decreases FOXC1 phosphorylation. (A) The nuclear extracts of HEK-293T cells transiently expressing the indicated FOXC1 mutants were treated with different combinations of CIP and sodium vanadate. After the treatments the recombinant proteins were detected via western immunoblot using an anti-myc antibody. The arrowhead indicates the position of the phosphorylated FOXC1. The sample loading control, endogenous LDH, was detected via immunoblot using an anti-LDH antibody (Chemicon). (B) The percentage of phosphorylated FOXC1 in untreated samples (CIP-, NaVO₃-) was estimated by densitometry of the upper band as indicated in Materials and Methods. Error bars correspond to the SD of three independent experiments carried out in triplicate. NT: non-transfected cells (negative control). The asterisk indicates statistical significance compared to the control: p<0.05 (*); One-way ANOVA.

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In addition, the predicted phosphorylation potential of the mutant Ser residue indicated that it was unphosphorylated (data not shown). These results suggest that p.I126S can alter the FOXC1 conformation, decreasing the phosphorylation of the whole protein.

**EMSA assay**

The DNA binding ability of the different mutations was evaluated using EMSA analysis. Competition experiments in which similar amounts of pre-bound protein-DNA complexes were challenged with increasing ratios of competitor DNA containing the FOXC1 binding site (ranging from 0.2 to 10 pmol, i.e., 1- to 50-fold excess), showed that both the intensity and the proportion of the protein-DNA complexes formed by the p.I126S and p.I126M (control) mutations, were significantly lower than those of the wild-type protein (Fig. 7A and B). Conversely, and although the intensity of the complexes corresponding to p.G447_G448insDG was higher than those of the normal protein (Fig. 7A), the DNA binding ability did not differ significantly from that of the wild-type protein.

(Fig. 6A, lanes 3–4, 7–8 and 11–12), indicating that the upper band corresponds to phosphorylated FOXC1. In addition, the predicted phosphorylation potential of the mutant Ser residue indicated that it was unphosphorylated (data not shown). These results suggest that p.I126S can alter the FOXC1 conformation, decreasing the phosphorylation of the whole protein.
from the wild-type (Fig. 7B). As expected, the two nonsense mutants failed to bind to the DNA because they lacked a complete forkhead domain (data not shown).

Subcellular localization
Fluorescence immunocytochemistry of the control recombinant wild-type FOXC1 localized the protein exclusively to the nucleus (Fig. 8, Wt). The two non-sense mutations were detected in both the nucleus and the cytoplasm (Fig. 8, p.Y47X and p.Q106X). Mutation p.I126S was also present in these two cell compartments. In contrast, the p.I126M control mutation was observed exclusively in the nucleus. p.G447_G448insDG also showed normal nuclear localization (Fig. 8).

Discussion
Glaucoma is a genetically and clinically heterogeneous disease with a poorly understood genetic basis. It has been proposed that dominant glaucoma may originate from goniodysgenesis with marked variation in the expressivity of dysgenesis, leading to the apparition of symptoms early or late in life [1]. Typical congenital glaucoma, which is inherited as a non-dominant trait, is also considered to be part of the anterior segment dysgenesis (ASD) group of disorders because it originates via the maldevelopment of structures in the anterior segment of the eye. On the converse, FOXC1 is a gene that regulates the development of the anterior segment of the eye, and it is known to be involved in several autosomal dominant eye defects associated with an increased risk for glaucoma. Its heterozygous loss-of-function in mice is associated with considerable variability in anterior segment abnormalities, similar to those present in patients with ARA and glaucoma [37]. Based on these facts, we explored the role of FOXC1 in dominant glaucoma. To this end, we carried out a comprehensive screening of the coding and regulatory regions (UTRs and the promoter) of the gene in twelve affected Spanish families with no known alterations in other glaucoma genes.

Pathogenicity of the FOXC1 mutations
We identified three hypermorphic (p.Y47X, p.Q106X and p.G447_G448insDG) and one hypomorphic (p.I126S) heterozygous FOXC1 variants in four probands, representing a significant proportion (33.3%) of the studied families. Only two of these mutations have previously been reported. p.Q106X was identified in a patient with an enucleated eye due to severe glaucoma and abnormalities of the anterior segment in the other eye [21]. Interestingly, p.Q106X-related mutations have been found to be associated with high phenotypic variability. For instance, p.Q120X was detected in patients with ARS and Peters’ anomaly in the same family [38], and p.Q123X has been identified in ARA patients and even in a normal subject [39].

It is accepted that FOXC1 mutations are pathogenic and function by reducing the levels of protein, inhibiting DNA binding and/or reducing the levels of FOXC1 transactivation [22, 40]. To the best of our knowledge, this is the first report of hypermorphic FOXC1 variants associated with dominant glaucoma, which agrees with previous studies showing a high glaucoma risk associated with increased gene dosage due to FOXC1 gene duplication [23, 41, 42]. The unexpected hyperactivity of the two truncated proteins is likely due to the expression of the isolated activation domain 1, which appears to act as a potent and autonomous transcription up-regulator despite the reduced stability of the mutant polypeptides. According to these results it has been described that expression of N-terminal FOXC1 residues 1–30 and 1–65 present transactivation levels that are equivalent to or higher than the full-length FOXC1, and it has been suggested that the N-terminal regions are sufficient to activate transcription [20]. Because the truncated proteins also failed to bind DNA, and pull-down experiments indicated that they did
FOXC1 Mutations in Dominant Glaucoma

not interact with wild-type FOXC1 (data not shown), we propose that the increased transactivation is a gain-of-function consisting in abnormal interactions with other transcription
factors. Our findings complements the haploinsufficiency mechanism, which has frequently been invoked to underlie the pathogenicity of FOXC1 non-sense alleles. The mislocalization of the truncated proteins (present in both the nucleus and the cytoplasm) is likely due to their small molecular size (< 50 kDa), which allows free diffusion through the nuclear pores. The remaining hypermorphic mutation, p.G447_G448insDG, did not differ from the wild-type protein in terms of the protein stability, DNA binding or subcellular localization. Nevertheless, its localization in a Gly-rich region, which is predicted to be unstructured (IDR4), may explain the observed hyperactivity. IDRs are known to lack a fixed tertiary structure and are present in different regulatory proteins, which include the transcription factors CREB [43] and p53 [44]. They participate in molecular interactions with other proteins or nucleic acids, and although they lack a defined structure when alone in solution, they undergo disorder-to-order transitions after binding to specific targets [45]. These data suggest that the increased transcription activity of p.G447_G448insDG may be also due to a gain-of-function that would result in altered binding to transcription factors, suggesting a possible role of the mutant amino acids in protein-protein interactions. p.I126S was hypomorphic and its decreased protein stability and DNA binding affinity were similar to that of the p.I126M control mutation. The reduction in DNA binding affinity may be due to structural alterations in the forkhead domain alpha-helix 3, which is predicted to play a role in major DNA groove recognition. Contrary to our results, p.I126M has been described to bind the FBE at near-wild-type FOXC1 levels [22]. This discrepancy may reflect different levels of excess unlabelled oligonucleotides used as competitors for FOXC1 DNA binding, which ranged from 1- to 50-fold in the present study and from 0.5- to 1.5-fold in a previous report. In contrast to p.I126M, p.I126S reduced FOXC1 phosphorylation and induced intracellular mislocalization of this protein. Because the amino acid residues involved in the p.I126M mutation are not phosphorylated, we postulate that a mutation-induced conformational change may interfere the interaction between kinases and FOXC1, leading to decreased FOXC1 phosphorylation. In accordance with our results, it has been described that I126 is critical for correct FOXC1 nuclear localization and transactivation, so that substitution of I126 for different amino acids leads to differentially disrupted nuclear localization and DNA binding [46]. Although the differences in protein phosphorylation and subcellular localization between these two mutations do not seem to correlate with the severity of the associated phenotypes, we should bear in mind that unidentified genetic modifiers may also contribute to the final phenotypic outcome.

Phenotypic variability and residual FOXC1 activity

Although dominant glaucoma was the most outstanding clinical feature in all of the mutation carriers, a variability in the age of onset was also observed. Moreover, one of the families (PCG73) also showed variable iris morphology, which ranged from thin stromal areas and posterior embrytoxon with no iris attachments (subjects II:2 and III:1), to corectopia associated with Axenfeld-Rieger anomaly (subject II:4). Mild iris alterations can be present in non-syndromic congenital glaucoma cases [47]. Previous reports have also identified a remarkable phenotypic variability associated with FOXC1 mutations [39, 48]. This clinical heterogeneity suggests an important role for modifier factors (genetic, environmental and/or stochastic) on the phenotypic outcomes. The FOXC1 associated phenotypic variability may also depend on the residual transcriptional activity associated with the individual genotypes. Based on this idea, we observed that none of the glaucoma-associated FOXC1 mutations expressed in the cell cultures showed a complete loss-of-function. On the contrary, the glaucoma associated transcriptional activity ranged from roughly 20% to 198% of the wild-type levels. Using an additive model, we estimated the FOXC1 transactivation ability associated with each glaucoma genotype as the mean activity of the two alleles and inferred two glaucoma associated activity...
intervals, which ranged from approximately 50% to 60% and from 130% to 150% compared to the wild-type FOXC1 genotype (Fig. 9). Based on this hypothesis we postulate that the variation in the residual transactivation between these narrow limits primarily results in goniodysgenesis and leads to dominant glaucoma as the most important phenotypic feature. Moreover, FOXC1 activity levels beyond the upper and lower thresholds would lead to severe ASD affecting multiple ocular and even non-ocular tissues. This model also suggests that the normal ocular development tolerates only small FOXC1 activity fluctuations (approximately ±35%). Consistent with this view, it has been proposed that FOXC1 haploinsufficiency (50% of the normal activity) underlies dominant glaucoma in ARA patients [35] and increased gene dosage due to heterozygous gene duplications (150% activity, representing the upper activity value of this model compatible with alterations limited to the eye), causes dominant inherited anterior-chamber defects associated with developmental glaucoma [49]. We also postulate an important influence of modifier genes, environmental factors and/or stochastic events on these transactivation...
thresholds. This hypothesis agrees with the proposed existence of critical activity thresholds for FOXC1 and the related protein PITX2 [42].

Conclusions
In summary, this study shows that FOXC1 sequence variations with moderate activity defects are associated with dominant glaucoma and remarkable phenotypic variability. In addition, this is the first study to identify hypermorphic variants of this gene and will contribute to a better understanding of the genetic basis of dominant glaucoma, as well as the function of FOXC1.

Supporting Information
S1 Fig. Localization of primers used in this study to analyze FOXC1. The arrows show the position of the different PCR primers. Amplicons are indicated by horizontal brackets and the numbers between parentheses correspond to amplicon length. CR: coding region. (TIF)

S2 Fig. Tricine-SDS-PAGE electrophoresis of the truncated FOXC1 mutations. To improve detection of the small molecular size mutants p.Y47X and p.Q106X, the nuclear extracts of HEK-293T cells transiently expressing these recombinant proteins were analyzed via Tricine-SDS-PAGE. The proteins were detected via western immunoblot using an anti-myc antibody. NT: non-transfected. (TIF)

S1 Materials and Methods. (DOCX)

S1 Table. The primer sequences and PCR conditions used for FOXC1 sequencing. (DOCX)

S2 Table. The primer sequences and PCR conditions used for FOXC1 site-directed mutagenesis. (DOCX)

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Author Contributions
Conceived and designed the experiments: JE JGF CMT. Performed the experiments: CMT FSS JDAA JJFF. Analyzed the data: CMT FSS JDAA JJFF LM CDMH FBK CA JGF JE. Contributed reagents/materials/analysis tools: LM CDMH FBK CA JGF JE. Wrote the paper: CMT JDAA JJFF CDMH CA JGF JE.
References

1. Jemdal T. Congenital glaucoma due to dominant goniodysgenesis. A new concept of the heredity of glaucoma. Am J Hum Genet. 1983; 35(4): 645–51. PMID: 6881141

2. Simha N, Verin P, Gauthier L. Congenital glaucoma of dominant autosomal transmission apropos of a family. Management. Bull Soc Ophthalmol Fr. 1989; 89(10): 1149–51. PMID: 2620399

3. Chakrabarti S, Kaur K, Rao KN, Mandal AK, Kaur I, Parkh RS, et al. The transcription factor gene FOXC1 exhibits a limited role in primary congenital glaucoma. Invest Ophthalmol Vis Sci. 2009; 50(1): 75–83. doi: 10.1167/iovs.08-2253 PMID: 18708620

4. Stolov I, Akarsu AN, Sarfarazi M. Identification of three different truncating mutations in cytochrome P4501B1 (CYP1B1) as the principal cause of primary congenital glaucoma (Buphthalmos) in families linked to the GLC3A locus on chromosome 2p21. Hum Mol Genet. 1997; 6(4): 641–7. PMID: 9097971

5. Ali M, McKibbin M, Booth A, Parry DA, Jain P, Riazuddin SA, et al. Null mutations in LTBP2 cause primary congenital glaucoma. Am J Hum Genet. 2009, 84(5): 664–71. doi: 10.1016/j.ajhg.2009.03.017 PMID: 19361779

6. Bagiyeva S, Marfany G, Gonzalez-Angulo O, Gonzalez-Duarte R. Mutational screening of CYP1B1 in Turkish PCG families and functional analyses of newly detected mutations. Mol Vis. 2007; 13: 1458–68. PMID: 17893647

7. Lopez-Garrido MP, Medina-Trillo C, Morales-Fernandez L, Garcia-Feijoo J, Martinez-de-la-Casa JM, Garcia-Anton M, et al. Null CYP1B1 genotypes in primary congenital and non-dominant juvenile glaucoma. Ophthalmology. 2013; 120(4): 716–23. doi: 10.1016/j.ophtha.2012.09.016 PMID: 23218183

8. Bejjani BA, Lewis RA, Tomey KF, Anderson KL, Dueker DK, Jabak M, et al. Mutations in CYP1B1, the gene for cytochrome P4501B1, are the predominant cause of primary congenital glaucoma in Saudi Arabia. Am J Hum Genet. 1998; 62(2): 325–33. PMID: 9463332

9. Chakrabarti S, Kaur K, Komatireddy S, Acharya M, Devi KR, Mukhopadhyay A, et al. Gln48His is the prevalent myocilin mutation in primary open angle and primary congenital glaucoma phenotypes in India. Mol Vis. 2005; 11: 111–3. PMID: 15729004

10. Kaur K, Reddy AB, Mukhopadhyay A, Mandal AK, Hasnain SE, Ray K, et al. Myocilin gene implicated in primary congenital glaucoma. Clin Genet. 2005; 67(4): 335–40. PMID: 15733270

11. Stone EM, Fingert JH, Alward WL, Nguyen TD, Polansky JR, Sunden SL, et al. Identification of a gene that causes primary open angle glaucoma. Science. 1997; 275(5300): 668–70. PMID: 9005853

12. Wiggs JL, Allingham RR, Vollrath D, Jones KH, De La PM, Kern J, et al. Prevalence of mutations in TIGR/Myocilin in patients with adult and juvenile primary open-angle glaucoma. Am J Hum Genet. 1998; 63(5): 1549–52. PMID: 9792882

13. Rezaie T, Child A, Hitchings R, Brice G, Miller L, Coca-Prados M, et al. Adult-onset primary open-angle glaucoma caused by mutations in optineurin. Science. 2002; 295(5557): 1077–9. PMID: 11834836

14. Pasutto F, Keller KE, Weisschuh N, Sticht H, Samples JR, Yang YF, et al. Variants in ASB10 are associated with open-angle glaucoma. Hum Mol Genet. 2012; 21(6): 1336–49. doi: 10.1093/hmg/ddr572 PMID: 22156576

15. Monem S, Spaeth G, DaSilva A, Popinchalk S, Iltichev E, Liebmann J, et al. Identification of a novel adult-onset primary open-angle glaucoma (POAG) gene on 5q22.1. Hum Mol Genet. 2005; 14(6): 725–33. PMID: 15677485

16. Nishimura DY, Swiderski RE, Alward WL, Seary CC, Patil SR, Bennet SR, et al. The forkhead transcription factor gene FKHL7 is responsible for glaucoma phenotypes which map to 6p25. Nat Genet. 1998; 19(2): 140–7. PMID: 9620769

17. Lines MA, Kozlowski K, Walter MA. Molecular genetics of Axenfeld-Rieger malformations. Hum Mol Genet. 2002; 11(10): 1177–84. PMID: 12015277

18. Kume T, Deng KY, Winfrey V, Gould DB, Walter MA, Hogan BL. The forkhead/winged helix gene Mf1 is disrupted in the pleiotropic mouse mutation congenital hydrocephalus. Cell. 1998; 93(6): 985–98. PMID: 9635428

19. Kidson SH, Kume T, Deng K, Winfrey V, Hogan BL. The forkhead/winged-helix gene, Mf1, is necessary for the normal development of the cornea and formation of the anterior chamber in the mouse eye. Dev Biol. 1999; 211(2): 306–22. PMID: 10395790

20. Berry FB, Saleem RA, Walter MA. FOXC1 transcriptional regulation is mediated by N- and C-terminal activation domains and contains a phosphorylated transcriptional inhibitory domain. J Biol Chem. 2002; 277(12): 10292–7. PMID: 11782474

21. D’Haene B, Meire F, Claerhout I, Kroses H, Plomp A, Arens YH, et al. Expanding the spectrum of FOXC1 and PITX2 mutations and copy number changes in patients with anterior segment malformations. Invest Ophthalmol Vis Sci. 2011; 52(1): 324–33. doi: 10.1167/iovs.10-5309 PMID: 20881294
22. Saleem RA, Banerjee-Basu S, Berry FB, Baxevanis AD, Walter MA. Analyses of the effects that disease-causing missense mutations have on the structure and function of the winged-helix protein FOXC1. Am J Hum Genet. 2001; 68(3): 627–41. PMID: 11179011

23. Lehmann OJ, Ebenezer ND, Jordan T, Fox M, Ocaka L, Payne A, et al. Chromosomal duplication involving the forkhead transcription factor gene FOXC1 causes iris hypoplasia and glaucoma. Am J Hum Genet. 2000; 67(5): 1129–35. PMID: 11007653

24. Strungaru MH, Dinu I, Walter MA. Genotype-phenotype correlations in Axenfeld-Rieger malformation and glaucoma patients with FOXC1 and PITX2 mutations. Invest Ophthalmol Vis Sci. 2007; 48(1): 228–37. PMID: 17197537

25. Lopez-Garrido MP, Sanchez-Sanchez F, Lopez-Martinez F, Aroca-Aguilar JD, Blanco-Marchite C, Coca-Prados M, et al. Heterozygous CYP1B1 gene mutations in Spanish patients with primary open-angle glaucoma. Mol Vis. 2005; 11: 748–55. PMID: 16862072

26. Aroca-Aguilar JD, Sanchez-Sanchez F, Ghosh S, Coca-Prados M, Escribano J. Myocilin mutations causing glaucoma inhibit the intracellular endoproteolytic cleavage of myocilin between amino acids Arg226 and Ile227. J Biol Chem. 2005; 280(22): 21043–51. PMID: 15795224

27. Sanchez-Sanchez F, Martinez-Redondo F, Aroca-Aguilar JD, Coca-Prados M, Escribano J. Characterization of the intracellular proteolytic cleavage of myocilin and identification of calpain II as a myocilin-processing protease. J Biol Chem. 2007; 282(38): 27810–24. PMID: 17650508

28. Hayashi H, Kume T. Forkhead transcription factors regulate expression of the chemokine receptor CXCR4 in endothelial cells and CXCL12-induced cell migration. Biochem Biophys Res Commun. 2008; 367(3): 584–9. doi: 10.1016/j.bbrc.2007.12.183 PMID: 18187037

29. Saleem RA, Banerjee-Basu S, Berry FB, Baxevanis AD, Walter MA. Structural and functional analyses of disease-causing missense mutations in the forkhead domain of FOXC1. Hum Mol Genet. 2003; 12(22): 2993–3005. PMID: 14501633

30. Saleem RA, Murphy TC, Liebmann JM, Walter MA. Identification and analysis of a novel mutation in the FOXC1 forkhead domain. Invest Ophthalmol Vis Sci. 2003; 44(11): 4608–12. PMID: 14578375

31. Campos-Mollo E, Lopez-Garrido MP, Blanco-Marchite C, Garcia-Feijoo J, Peralta J, Belmonte-Martínez J, et al. CYP1B1 mutations in Spanish patients with primary congenital glaucoma: phenotypic and functional variability. Mol Vis. 2009; 15: 417–31. PMID: 19234632

32. Berry FB, Lines MA, Oas JM, Fozz T, Underhill DA, Gage PJ, et al. Functional interactions between FOXC1 and PITX2 underlie the sensitivity to FOXC1 gene dose in Axenfeld-Rieger syndrome and anterior segment dysgenesis. Hum Mol Genet. 2006; 15(6): 905–19. PMID: 16449236

33. Schagger H, von Jagow G. Tricine-sodium dodecyl sulfate-polyacrylamide gel electrophoresis for the separation of proteins in the range from 1 to 100 kDa. Anal Biochem. 1987; 166(2): 368–79. PMID: 2449095

34. Romero-Calvo I, Ocon B, Martinez-Moya P, Suarez MD, Zarzuelo A, Martinez-Augustín O, et al. Reversible Ponceau staining as a loading control alternative to actin in Western blots. Anal Biochem. 2010; 401(2): 318–20. doi: 10.1016/j.ab.2010.02.036 PMID: 20206115

35. Mears AJ, Jordan T, Mirzayans F, Dubois S, Kume T, Parlee M, et al. Mutations of the forkhead/winged-helix gene, FKHL7, in patients with Axenfeld-Rieger anomaly. Am J Hum Genet. 1998; 63(5): 1316–28. PMID: 9792859

36. Papadopoulos M, Brandt JD, Sugiyama K, Khaw PT, Chu J, Law S, et al. Establisng the diagnosis and determining glaucoma progression. In: Weinreb RN, Graniewski AL, Papadopoulos M, Grigg J, Freedman S, editors. Childhood Glaucoma. Ámsterdam, The Netherlands: Kluger Publications; 2013. p. 15–41. doi: 10.1186/1687-9856-2013-15 PMID: 24025597

37. Smith RS, Zabaleta A, Kume T, Savinova OV, Kidson SH, Martin JE, et al. Haploinsufficiency of the transcription factors FOXC1 and FOXC2 results in aberrant ocular development. Hum Mol Genet. 2004; 13(19): 2429–37. doi:10.1093/hmg/ddh292 PMID: 15177473
41. Lehmann OJ, Ebenezer ND, Ekong R, Ocakes L, Mungall AJ, Fraser S, et al. Ocular developmental abnormalities and glaucoma associated with interstitial 6p25 duplications and deletions. Invest Ophthalmol Vis Sci. 2002; 43(6): 1843–9. PMID: 12036988

42. Walter MA. PITs and FOXes in ocular genetics: the Cogan lecture. Invest Ophthalmol Vis Sci. 2003; 44(4): 1402–5. PMID: 12657570

43. Sugase K, Dyson HJ, Wright PE. Mechanism of coupled folding and binding of an intrinsically disordered protein. Nature. 2007; 447(7147): 1021–5. PMID: 17522630

44. Tidow H, Melero R, Mylonas E, Freund SM, Grossmann JG, Carazo JM, et al. Quaternary structures of tumor suppressor p53 and a specific p53 DNA complex. Proc Natl Acad Sci USA. 2007; 104(30): 12324–9. PMID: 17620598

45. Uversky VN, Gillespie JR, Fink AL. Why are “natively unfolded” proteins unstructured under physiologic conditions? Proteins. 2000; 41(3): 415–27. PMID: 11025552

46. Saleem RA, Banerjee-Basu S, Murphy TC, Baxevanis A, Walter MA. Essential structural and functional determinants within the forkhead domain of FOXC1. Nucleic Acids Res. 2004; 32(14): 4182–93. PMID: 15299087

47. Perry LP, Jakobiec FA, Zakka FR, Walton DS. Newborn primary congenital glaucoma: histopathologic features of the anterior chamber filtration angle. J AAPOS. 2012; 16(6): 565–8. doi: 10.1016/j.jaapos.2012.06.012 PMID: 23158552

48. Perveen R, Lloyd IC, Clayton-Smith J, Churchill A, van Heyningen V, Hanson I, et al. Phenotypic variability and asymmetry of Rieger syndrome associated with PITX2 mutations. Invest Ophthalmol Vis Sci. 2000; 41(9): 2456–60. PMID: 10937553

49. Nishimura DY, Searby CC, Alward WL, Walton D, Craig JE, Mackey DA, et al. A spectrum of FOXC1 mutations suggests gene dosage as a mechanism for developmental defects of the anterior chamber of the eye. Am J Hum Genet. 2001; 68(2): 364–72. PMID: 11170889

50. Linding R, Jensen LJ, Diella F, Bork P, Gibson TJ, Russell RB. Protein disorder prediction: implications for structural proteomics. Structure. 2003; 11(11): 1453–9. PMID: 14604535

51. Linding R, Russell RB, Neduva V, Gibson TJ. GlobPlot: Exploring protein sequences for globularity and disorder. Nucleic Acids Res. 2003; 31(13): 3701–8. PMID: 12824398

52. Thompson JD, Higgins DG, Gibson TJ. CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res. 1994; 22(22): 4673–80. PMID: 7984417