Stargardt Phenotype Associated With Two *ELOVL4* Promoter Variants and *ELOVL4* Downregulation: New Possible Perspective to Etiopathogenesis?

Luigi Donato,¹⁻³ Concetta Scimone,^{2,3} Carmela Rinaldi,³ Pasquale Aragona,³ Silvana Briuglia,³ Angela D'Ascola,³ Rosalia D'Angelo,³ and Antonina Sidoti^{2,3}

¹Department of Chemical, Biological, Pharmaceutical and Environmental Sciences, University of Messina, Messina, Italy ²Department of Cutting-Edge Medicine and Therapies, Biomolecular Strategies and Neuroscience, Section of Neuroscience-Applied Molecular Genetics and Predictive Medicine, Istituto Euro Mediterraneo di Scienza e Tecnologia (I.E.ME.S.T.), Palermo, Italy ³Department of Biomedical and Dental Sciences and Morphofunctional Imaging, University of Messina, Messina, Italy

Correspondence: Rosalia D'Angelo, Department of Biomedical and Dental Sciences and Morphofunctional Imaging, Division of Molecular Genetics and Preventive Medicine, University of Messina, via C. Valeria 1, I-98125 Messina, Italy; rdangelo@unime.it.

Submitted: September 11, 2017 Accepted: January 7, 2018

Citation: Donato L, Scimone C, Rinaldi C, et al. Stargardt phenotype associated with two *ELOVL4* promoter variants and *ELOVL4* downregulation: new possible perspective to etiopathogenesis? *Invest Ophtbalmol Vis Scl.* 2018;59:843–857. https://doi.org/ 10.1167/iovs.17-22962 **PURPOSE.** Stargardt disease (STGD) is the most common form of inherited juvenile macular degeneration. It is inherited as autosomal recessive trait (STGD1), although STGD3 and STGD4 are inherited as autosomal dominant inheritance pattern. STGD3 is caused by mutations in the elongation of very long-chain fatty acids-like 4 (*ELOVL4*) gene encoding for a very long-chain fatty acids-like 4 (*ELOVL4*) gene encoding for a very long-chain fatty acids-like 4 (*ELOVL4*) gene encoding for a very long-chain fatty acids elongasic reticulum. STGD occurs due to altered synthesis of very long-chain polyunsaturated fatty acids (VLC-PUFA). Our work investigates the role of two variants in the *ELOVL4* gene promoter region, c.-236 C>T (rs240307) and c.-90 G>C (rs62407622), identified in a patient with STGD in transconfiguration.

METHODS. Their effects on *ELOVL4* expression were examined by Dual-Luciferase Reporter assay.

RESULTS. rs62407622 and rs240307 variants caused 14% and 18% of expression reduction, respectively, compared with wild-type promoter. A very strong decreased gene expression was caused by coexistence of both variants.

CONCLUSIONS. A highly reduced activity of the *ELOVL4* promoter was registered due to combination of two variants. Decrease of *ELOVL4* enzymatic activity could lead to a deficiency of VLC-PUFA, essential components for rods function and longevity, which are among the parameters involved in the etiopathogenesis of STGD.

Keywords: Stargardt disease, macular degeneration, retinal degeneration, photoreceptors, VLC-PUFA

S targardt disease (STGD) is a form of retinal dystrophy usually characterized by a progressive loss of central vision associated with irregular macular and perimacular yellow-white fundus flecks, and a so-called "beaten bronze" atrophic central macular lesion. Worldwide prevalence of STGD is estimated between 1/8000 and 1/10,000.¹

Typical disease onset does not exceed the twentieth year of life, although symptoms can also appear during adulthood and as late as the seventh decade.² Although disease severity and progression varies widely, STGD is usually characterized by loss of visual acuity, followed by wavy vision, blind spots, blurriness, impaired color vision, photophobia, and difficulty adapting in the dark.³

Today, three forms of STGD are known: STGD1, STGD3, and STGD4.⁴ STGD3 (Online Mendelian Inheritance in Man [OMIM] #600110) is a rare dominant form due to mutations in the elongation of very long-chain fatty acids-like 4 (*ELOVL4*) gene on chromosome 6q16.^{5–8} *ELOVL4* plays a fundamental role in the synthesis of very long-chain polyunsaturated fatty acids (VLC-PUFA).^{9,10} VLC-PUFA make up a considerable part of phosphatidylcholine (PC) in the outer segment of both cell types of photoreceptors, suggesting a relevant role in the

correct folding of disk rim and in cones and rods membrane fluidity.¹¹ These functions, together with a close interaction with rhodopsin, strongly point to the possible involvement of *ELOVL4* in phototransduction.¹² Furthermore, recently, VLC-PUFA have also been found in conventional synapses and retina ribbon, probably being incorporated into vesicles containing glutamate, in rods terminals.¹³

So far only nine *ELOVL4* variants are known: six single nucleotide and three indel/del mutations. Among these, four variants are in exon 6 and associated with the STGD3 form.¹⁴ According to this data, it would appear that Elovl4 truncated protein loses the endoplasmic reticulum retention signal (KXKXX) and is mislocalized from the site of synthesis of VLC-PUFA. This condition leads to retina degeneration, due to: (1) production of toxic3-keto-acylintermediates that imply cell death, and (2) reduced levels of VLC-PUFA and mislocalization of mutant protein, along with cellular stress, causing impairment of important cellular functions.⁵ It cannot be excluded that, in these situation, several chaperons, like HSP90, could help to solve the problem, as in other ocular diseases.¹⁵

Copyright 2018 The Authors iovs.arvojournals.org | ISSN: 1552-5783

Investigative Ophthalmology & Visual Science

843

Here, we report the case of a patient with dominant STGD in which we identified two *ELOVL4* promoter variants, c.-236 C>T (rs240307) and c.-90 G>C (rs62407622).

The effects of the single c.-90 G>C and c.-236 C>T variants, as well as two variants together ones (c.-90 G>C and c.-236 C>T) on gene expression and, consequently, on the onset of disease, were examined.

MATERIALS AND METHODS

Clinical Data

The proband, a 42-year-old Caucasian man, came to our attention with a diagnosis of STGD, showing visual problems since young age. His visual acuity was 1.6/10 in the right eye and 2/10 in the left eye; his peripheral visual field was well represented, while the central one was almost absent. Moreover, he also showed an initial loss of color vision, photophobia, and a slow dark adaptation. Diagnosis was the result of the following evaluations: fundus analysis, fundus autofluorescence (FAF), infrared reflectance imaging (IR), optical coherent tomography (OCT), visual field (VF), International Society for Clinical Electrophysiology of Vision (ISCEV) ERG, and pattern electroretinogram (PERG). Fundus examination revealed bilateral anatrophic, rounded maculopathy with sharp edges, surrounded by pisciform flecks, confirmed by IR and FAF (Fig. 1). Furthermore, FAF showed mottled areas of hyperautofluorescence and hypoautofluorescence, corresponding to areas of lipofuscin accumulation and RPE atrophy, respectively. ERG revealed a generalized rods dysfunction with cones involvement (photopic and scotopic hypovolted ERG), with a delay in visual response (PERG with hypovolted P50 wave and increased latency; BiomedicaMangoni, Pisa, Italy; Fig. 2). VF showed central scotoma correlating with outer retinal subfoveal atrophy observed on FAF and OCT (Fig. 3). In details, OCT highlighted disruption of both inner and outer photoreceptor segment layers, combined with the loss of the inner segment-outer segment junction and thinning of other retina layers.

The patient's family, composed of father and mother, was evaluated by the same clinical and instrumental analyses and resulted healthy. Both parents did not manifest bilateral central visual loss, photophobia, color vision abnormalities, central scotomas, or slow dark adaptation. Moreover, they showed a visual acuity of 20/20, a normal visual field, and a clean fundus.

We screened all three known STGD causative genes (*ABCA4*, *ELOVL4*, and *PROM1*), and we found no associated or causative variants (HGMD Professional was the most important and updated database we considered; Qiagen Aarhus Prismet, Aarhus, Denmark), except those we analyzed in this paper.

In order to evaluate the variants effects on *ELOVL4* expression, we performed a genetic analysis of the gene promoter through PCR and Sanger sequencing, followed by an in silico prediction and the functional Dual-Luciferase Reporter assay. The latter was essential to experimentally confirm the previously generated data.

Genotyping

DNA was extracted from leukocytes by using standard protocols. Amplification of regulatory regions of *ELOVL4* gene was performed using primer pairs designed according to the published nucleotide sequence of GenBank (accession no. NG_009108.1; available upon request).

The PCR mix was prepared by adding 8 μ g of genomic DNA to 50 μ L reaction mixture containing a 0.2 μ m concentration of

IOVS | February 2018 | Vol. 59 | No. 2 | 844

each primer and 1 U MyTaq polymerase (Bioline, Aurogene Srl, Rome, Italy). PCR was carried out in the thermal cycler (Gene Amp PCR System 2700; PE Applied Biosystems, Foster City, CA, USA) under the following conditions: denaturation at 95°C for 15 seconds, annealing at 49.5°C for 15 seconds, and extension at 72°C for 10 seconds for 35 cycles, after an initial 1 minute denaturation at 95°C.

PCR products were sequenced by direct sequencing, using BigDye Terminator (ThermoFisher Scientific, Life Technologies, Monza, Italy) and Applied Biosystems 3500 Genetic Analyzer. The nucleotide number relative to variants identified in the promoter region of 805 bp was indicated in respect to the transcriptional start site of the reference sequence reported by the National Center for Biotechnology Information (NCBI) database. To name two polymorphisms we relied on Human Genome Variation Society (HGSV) nomenclature. Therefore, c.-90G> C (rs62407622) and c.-236C> T (rs240307) indicated nucleotide substitutions at RNA level.

In Silico Analysis

In silico analysis was performed on the *ELOVL4* promoter using two transcription factors (TF) prediction tools, individually and in pairs: BioBase TRANSFACTM Professional¹⁶ and Alggen PROMO.¹⁷ These tools were used to identify potential TF binding sites in the region where the variants in the *ELOVL4* promoter were found. TRANSFAC was set to use the profile matrix for vertebrates, with a cutoff to minimize false positives (minFP). This is defined as the score that gives 1% of hits in the used sequences relative to the number of hits received at the minimum false negative (minFN) cutoff (the score at which at least 90% of the positive test set are recognized, i.e., it equals a false negative rate of 10%). The false positive rate is estimated by applying the Match algorithm (BIOBASE Gmbh, Halchtersche, Wolfenbüttel, Germany) to upstream sequences.

PROMO, instead, involves the dissimilarity threshold; a parameter that controls how similar a sequence must be to the matrix to be reported as a hit. It was set at 15% (85% similarity). Random expectation (RE) gives the number of expected occurrences of the match, in a random sequence of the same length as the query sequence, according to the dissimilarity index. Two models are considered: (1) equiprobability for the four nucleotides (RE equally), and (2) estimate the nucleotide probability as the nucleotide frequencies in the query sequence (RE query).

Furthermore, Cytoscape software (The Cytoscape Consortium, New York, NY, USA) and its MCODE plug-in were used to analyze pathways between involved TFs, in order to predict possible interactions among them.

Cell Culture

U373 MG (human, Caucasian, glioblastoma-astrocytoma) cells (Sigma-Aldrich, Milan, Italy) were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 U/mL penicillin, and 1 mg/mL ampicillin (Lonza, Amboise, France) at 37° C in a water-saturated atmosphere with 5% CO₂. The essential feature that led us to choose this cell line was the acquisition of infinite growth potential, which set the stage for multiplication of genetic variants with an ever increasing fitness for proliferation and spread.

We used a glioblastoma cells line to perform another dual luciferase assay, involving cerebral cavernous malformations.¹⁸ This cell line can be useful because the retina has a nervous derivation as glia. Despite this, it is not possible to exclude that the whole transcription factors set in glioblastoma cells could be quite different from that in retina cells. In order to clarify



FIGURE 1. Infrared images (A, B), FAF (C, D), and fundus photograph (E, F), for the Proband compared with overt STDG3 (G-I) and control (J, K). Both *right* (C) and *left* (D) eyes show absence of FAF in the fovea, corresponding to geographic atrophy. Moreover, (A, B) evidence a central, bright, hyperreflective area, which corresponded to the atrophic macular lesion, surrounded by a darker, hyporeflective zone with distinct borders. This area is consistent with a zone where there is thinning and hypopigmentation of the RPE. This appearance is similar to established STDG3 in another patient, as evidenced in fundus photograph (G), fluorescein angiogram (H), and FAF (I). The fundus photograph and FAF of a healthy control are shown in (J, K).

this point, we address some supporting evidence: almost all TFs coming from our in silico analysis result expressed both in glioblastoma cells and retina cells, as evidenced by DPRP database,¹⁹ based on expression and ChIP-Seq experimental data, and TiGER.²⁰ Moreover, description of common TFs was reported among others also by Rheinbay et al.²¹ and Mysickova et al.²²

Construction of the Reporter Gene Plasmids

In order to highlight if the variants could affect *ELOVL4* expression, the effects of two combined genotypes, c.-236T/c.-90C (T-C) and c.-236C/c.-90C (C-C), on promoter activity were studied. An 805 bp promoter sequence was amplified by PCR using genomic DNA from the proband and one donor, selectively carrying each haplotype, using the following primers:

forward: 5'-AGATCTACATGCACTTCTCTTGTC-3' and reverse: 5'-AAGCTTCACTACGTTTAGGACAC-3' under these conditions: 1 cycle of 95°C for 1 minute; 35 cycles of 95°C for 15 seconds, 49.5°C for 15 seconds, and 72°C for 10 seconds; and 1 cycle of 72°C for 7 minute.

Each PCR product, as well as pGL4.10 (*luc2*) was digested by *BgIII* and *HindIII* (Promega Italia, Milan, Italy) and then purified (PureLink PCR Purification Kit; ThermoFisher Scientific, Life Technologies). Each promoter and pGL4.10 (*luc2*), digested and purified, was incubated overnight in appropriate concentrations, in order to execute ligation. The efficiency of promoter insertion upstream of the luciferase gene, cloned into the pGL4.10 (*luc2*), was verified by agarose gel electrophoresis.



FIGURE 1. Continued.



FIGURE 2. Proband's ERG and PERG, compared with healthy control. PERG (**A**) The voltage of the P50 wave was abnormally decreased and the latency was increased. The scotopic (**C**) and photopic (**E**) ERGs showed abnormally low voltage. The combination of this data suggests an impairment of both rods and cones functions, compared with PERG (**B**), scotopic (**D**), and photopic (**F**) reference ERGs. The signal was amplified (gain 50,000), filtered (band pass, 1–100 Hz) and averaged with automatic rejection of artifacts by a BM 6000 unit. Analysis time was 250 msec. In healthy subjects, these peaks have the following implicit times: 35, 50, and 95 msec (N35, P50, N95).



FIGURE 3. OCT of proband (**A**, **B**), compared with that of a healthy control (**C**) and overt STDG3 patient (**D**). Proband's right eye (**A**) depicts the extent of the transverse loss of the junction between the inner and outer segment of the photoreceptors in the foveal region. Furthermore, his left eye (**B**) showed abnormal pigmentation in the RPE layer, due to macular degeneration, as notable in the STDG3 patient (**D**). The integrity of all retina layers in a healthy subject is highlighted in (**C**).

FIGURE 4. Family tree genotypes and sequencing peaks for *ELOVL4* (6q14.1). Mutated condition, due to present polymorphisms, is *underlined*. *Black arrow* indicates the proband.

Novel constructs were subcloned into *Escherichia coli* Top 10 cells (Life Technologies) and single colonies were miniprep. The correct sequence of all the clones was verified by DNA sequencing, using the Sanger method, and then selected for transient transfection.

Transient Transfection and Promoter Assays

Cells were first seeded in 96-well culture plates at a density of 2 $\times 10^4$ cells per well. Then, a transient transfection was performed with following modality: (1) in 1/4 of wells (16), cells were cotransfected with 0.05 µg of the pGL4.10 [*luc2*] promoter construct containing the only c.-90 G>C variant, and with 0.05 µg of the pGL4.10 [*luc2*] promoter construct containing the *ELOVL4* wild-type promoter; (2) in a second 1/4 of wells (16), cells were cotransfected with 0.1 µg of the pGL4.10 [*luc2*] promoter construct containing only the c.-236 C>T variant, and with 0.05 µg of the pGL4.10 [*luc2*] promoter construct containing the *ELOVL4* wild-type promoter; (3) in another 1/4 of wells (16), cells were cotransfected with 0.05 µg of two pGL4.10 [*luc2*] promoter constructs containing, respectively, the c.-90 G>C and the c.-236 C>T variants; and

(4) in the final quarter of wells (16), cells were transfected with 0.1 µg of the pGL4.10 [luc2] promoter construct containing the ELOVL4 wild-type promoter. Finally, the remaining 32 wells were filled with not transfected cells (16) and with the luciferase substrate only (16). In each well, besides cells, the mixture included 0.2 µL of Lipofectamine 3000 Reagent (ThermoFisher Scientific, Waltham, MA, USA) and 0.2 µL of P3000 Reagent (ThermoFisher Scientific), in a serum-free medium and then incubated for 24 hours at 37°C in a humidified atmosphere of 5% CO₂ in air. After incubation, cells were washed twice with PBS and lysed by Passive Lysis Buffer (Promega). Luciferase activity was measured using Dual-Luciferase assay kit (Promega) and GloMax-Luminometer (Promega). Reporter construct activity was normalized by comparison with activity from the Renilla luciferase construct. Luciferase activities are representative of at least six independent experiments, with each construct tested 16 times per experiment.

Population Screening

Analyzed variants were screened in 500 unrelated healthy donors born and living in Messina for at least two generations, constituting a heterogeneous group for age and sex, in order to assess their frequency in the same geographic site population in which the patients belonged.

Statistical Analysis

All data analyses were performed using the IBM SPSS 24 software for Macintosh (IBM SPSS Statistics for Macintosh; IBM Corp, Armonk, NY, USA). A 1-way ANOVA was performed to compare between the sample groups. All P values were Bonferroni's corrected and considered significant if P < 0.05.

Ethical Statements

The study followed the tenets of the Declaration of Helsinki and was approved by the Scientific Ethics Committee of the Azienda Ospedaliera Universitaria-Policlinico "G. Martini" Messina. All family members and controls signed informed consent after explanation of the nature and possible consequences of the study.

RESULTS

Genotyping of ELOVL4 promoter in the whole family components highlighted that the proband's mother and father present an alternate heterozygosity for each variant in exam (Fig. 4), indicating that the probandinherited both variants in trans. The predictive analysis of the proband's ELOVL4 promoter by TRANSFAC Professional evidenced the loss of one group of TF binding sites (ETF, ZF5, E2F-6, FBI-I, HDAC2, and TAFII250) in the double heterozygous genotype, due to the presence of c.-90 G>C for most of them. The exception was represented by FBI-I and the complement 666 through 679 binding sites for TAFII250, whose loss was attributable to c.-236 C>T. A second group analysis, resulting from the combined predictions of TRANSFAC and PROMO, revealed the appearance of new possible binding sites of different TFs (CPB, BCL6B secondary motif, Spi-B, Pax-4, RXR-alpha, GKLF, POLR3A, TFII-I, Pax-5, p53, SP1, and GR-alpha), determined by c.-236 C>T for Spi-B, Pax-4 and RXR-alpha, and by c.-90 G>C for others (for further details see Table 1). This data suggest a probable transcription variation, due to the altered balance of TF binding properties. Furthermore, it is important to understand the relationship between the analyzed TFs, and how each one could influence the others. This was examined by Cytoscape pathway analysis, along with its MCODE plug-in, from which arose a 4-cluster division that highlighted a relevant network involving most of the TFs in exam (Table 2; Fig. 5). These data were confirmed by Dual-Luciferase Reporter Assay, involving the proband's ELOVL4 promoter, compared in its wild form, versus both variants and c.-90 G>C and c.-236 C>T only samples. Results showed an expression reduction of approximately 14% in the c.-90 G>C sample and of approximately 18% in the c.-236 C>T one (compared with a healthy control), but a strong decrease ($\sim 97\%$) arose from the promoter carrying the combination of above variants (Fig. 6). The 1-way ANOVA, after Bonferroni's correction, confirmed the statistical significance of analysis (P < 0.05). Multiple comparison details are listed in Supplementary Materials.

These results take a particular value, considering that both analyzed variants showed a very low frequency distribution in Messina healthy population (c.-90 G>C: G frequency = 0.94, C frequency = 0.06; c. 236 C>T: C frequency = 0.95%, T frequency = 0.05), in contrast to what reported in the European population (in the public domain, http://www.en sembl.org/Homo_sapiens/Variation/Population?db=core;r=6: 79946869-79947869;v=rs62407622;vdb=variation;vf=12792

675; in the public domain, http://www.ensembl.org/Homo_sapiens/Variation/Population?db=core;r=6:79947015-7994801 5;v=rs240307;vdb=variation;vf=129670).

DISCUSSION

ELOVL4 encodes a member of the elongase family, expressed in retina, brain, skin, and sperm, involved in the elongation of very long-chain fatty acids.¹⁰ Although little is known about the role of this protein, data report that the contribution of the enzyme is to be found in the initial rate of VLC-PUFA production and condensation reactions between a fatty acyl-CoA and malonyl-CoA.⁵ The role of VLC- PUFA is fundamental. The most reliable hypothesis argues that the VLC-PUFA acyl chain may cover the entire bilayer, representing a flexible hinge at the rim site where the curvature of photoreceptor disk membranes is the greatest.9 At rim level, alteration of ELOVL4 could impair the turnover of photoreceptor disk membranes, due to a modified balance of fatty acid precursors. The direct consequence of this variation leads to an abnormal accumulation of lipofuscin granules, observed in the RPE of mutant retinas that may impair retinoic acid trafficking between RPE and photoreceptors.²³

Recent experiments have shown a reduction in rods ERG oscillatory potentials and scotopic threshold responses in *ELOVL4* KO mice, and presented biomorphologic evidence that the ERG changes are correlated with reduced VLC-PUFA and synaptic architecture. It may affect vesicle tethering or recycling pathways, as well as glutamate release mechanisms, due to VLC-PUFA interaction with synaptic proteins that mediate endo/exocytic activity or that were localized to the synaptic ribbon in photoreceptor terminals.¹³

It is known that *ELOVL4* mutations, alone or with *PROM1* mutations, could cause enzyme activity loss and, subsequently, the onset of the dominant form of STGD.²⁴

Analysis of *ELOVL4* gene sequence in our patient affected by STGD permitted us to identify two variants, transconfigured on the gene promoter. We demonstrated that the coexistence of two variants determined the down regulation of gene transcription. To be precise, the Dual-Luciferase Reporter assay highlighted the down regulation of *ELOVL4* transcription by 97% in the patient's sample (c.-90 G>C and c.-236 C>T). The possible protein elonging activity loss could lead to several consequences:

- 1. Compromising the integrity of photoreceptor membrane compartments, such as Golgi, or even the retinal pigment epithelium²⁵;
- 2. Causing a corresponding reduction of rhodopsin levels within outer segment disk membranes, or the production of abnormal hetero-oligomers with its membrane proteins, which may lead to alterations in membrane ultrastructure or biochemistry²⁶; and
- 3. Altering VLC-PUFA direct signaling and possible alteration of the rim site where the curvature of photoreceptor disk membranes is greater or the greatest. It was proposed that lipid molecules, such as docosahexaenoic (DHA), eicosapentaenoic, and arachidonic acids could activate specific receptors or modulate transient receptor potential cation channel activity. The latter task is enforced by the presence of VLC-PUFAs also in ribbon synapses, as well as the smaller conventional synapses in the retina.¹³

Results showed that many TF binding sites were altered in the *ELOVL4* promoter, both for activators (ETF,²⁷ FBI-I,²⁸ HDAC2,²⁹⁻³² and TAFII250,^{33,34} Spi-B,³⁵ Pax-4,^{36,37} POL-R3A,^{38,39} TFII-I,^{40,41} PAX-5,^{42,43} TP53,⁴⁴⁻⁴⁶ SP1,⁴⁷⁻⁴⁹ and GR-

HereMatchCoreMatchCoreMatchCoreMatchCoreC.236C.236C.236C.236FF810.817minsCGCGCG0.840.8571.966.5C.37Reference (PMID)FF810.817minsCGCGCG0.840.8570.857V.8715,00777777FF810.817minsCGCGCGC0.840.857V.8715,00777777FH60.667minsCGCGCGC0.84V.8714,057777777FH/R210581.812pinsCGCGCGC0.84V.8714,057777777FH/R210581.812pinsCGCGCGC0.84V.8714,0577777777FH/R210565.670pinsCGCGCGC0.7580.840V.8714,10577772605066FHpinsCGCGCGC0.7580.840V.8714,10577726050667FLpinsCGCGCGC0.7580.840V.8714,1057772605066FLpinsCGCGCGC0.7580.840V.8714,1057772605066FLpinsCGCGCGC0.758V.8714,105777260506626050FLpinsCGCGCGC10.959V.87						BioBase	Transfac									
	Factor Name	Region	Strand	Match Sequence	Core Similarity	Matrix Similarity	Matrix ID	ΨŢ	c236 C>T c90 G>C	c90 G>C	c236 C>T	Reference (PMID)				
	ETF	807813	minus	CCGCCgc	1	1	V\$ETF_Q6	>			`	2768275				
	ZF5	810817	minus	ccgCTCGC	0.844	0.896	V\$ZF5_01	>			>	17714511				
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	E2F-6	811819	minus	cgctCGCCC	0.69	0.787	$V \pm E2F6_02$	>			>	9501179				
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	FBI-I	660668	minus	gGGCCTcct	0.625	0.79	V\$ZBTB7A_02	>		>		19853566; 15917220				
	HDAC2	811819	plus	CGCTCgccc	0.829	0.82	V\$HDAC2_04	>			>	26129908; 23940731; 23696608				
	TAFII250	666679	minus	CCTCCacctcctcg	0.807	0.84	V \$TAF1_05	>		>		18171927				
		804812	plus	cGGCCGccg	0.795	0.846	V \$TAF1_06	>			>					
		804812	minus	cggCCGCCg	0.798	0.849	V \$TAF1_06	>			>					
CPBP 809.815 plus cccccct 1 1 VSCPBP_Q6 7 7 24906800 178936461 16431354 BCL6B 803.818 plus crtcrCcc 1 0.979 V38DL6B.04 7 7 24906860 178936461 16431354 BCL6B 803.818 plus crtcrCcc 1 0.979 V38DL6B.04 7 7 24906860 178935461 16431354 BX4 664.674 plus crtrcrCcc 1 0.918 V38A4_02 7 7 24905860 179355661 16531354 RXRapha 664.674 plus 0.972 0.982 V5RAL_10 7 7 26310877 19475627 19520230 GKLF 810.822 minus ccCCCCgccg 0.994 0.962 V5RAL_10 7 7 2702452 1931571 RXRA_D stord stord 0.962 V5RAL_10 7 7 2702462 19176527 19315577 193176527 1931765		806814	plus	GCCGCcgct	1	0.984	V \$TAF1_07	>			>					
	CPBP	809815	plus	GCCCCtc	1	1	V\$CPBP_Q6		`	>		24906860; 17893646; 16431954				
SpiB 665.670 plus TTCTC 1 V \$\$NX4, Q2 V	BCL6B	803818	plus	ccggcCGCCCctcgcc	1	0.979	V\$BCL6B_04		`	>		23057762				
	Spi-B	665670	plus	TTCTCc	1	1	V\$SPIB_Q3		`		>	1406622				
RXRalpta 662.668 minus gcCTTCT 0.972 0.982 VRXR_10$ \prime Z <t< td=""><td>Pax-4</td><td>664674</td><td>plus</td><td>cttctCCACCt</td><td>1</td><td>0.918</td><td>V\$PAX4_Q2</td><td></td><td></td><td></td><td>></td><td>19843539; 19012751</td></t<>	Pax-4	664674	plus	cttctCCACCt	1	0.918	V\$PAX4_Q2				>	19843539; 19012751				
GKLF 810.822 minus cCCTCgccccg 0.989 V\$GKLF_Q3_01 t t 2702622 POLR3A 812.824 minus ccCGCCccgcg 0.944 0.962 V\$POLR3A_01 t 2702622 POLR3A 812.824 minus ccCGCCccgcg 0.944 0.962 V\$POLR3A_01 t 2702622 PAL5 Alleen Match 0.962 0.910 t t 2702622 Fator Name Region Strand Scoucc 0.944 0.962 t t t 2702622 Fator Name Region Strand Match t t t t t Fator Name Region Strand Match RE Query t	RXRalpha	662668	minus	gcCTTCT	0.972	0.982	V\$RXRA_10		`		>	26310807; 19427305; 10520230				
	GKLF	810822	minus	cCCTCgccccgt	0.989	0.989	V\$GKLF_Q3_01		`	>		27022622				
Algene PROMOAuthMatchFactor NameRegionStrandSequenceDissimilarityRE quallyRE QueryWTC236 C.>TC30C356TFIH664.669plusTTCTCC11.3378881.245121.92792C90 G.>CC.>TRE freence (PMID)TFIH664.669plusTTCTCC11.3378881.245121.9483C90 G.>CC.>TN17239664TP33805.811plusGCCGCCCC9.5521050.747071.9483YN22278416TP33805.811plusGCCGCCCCC5.184980.311280.91106YY22778416SP1803.812plusCGGCGCCCC3.1707070.017510.09628YYY26439195; 25407019SP1811.815plusCCTCG8.2815683.984385.91804YY2211818SP1811.815plusCCTCG8.2815683.984385.91804YY2211818	POLR3A	812824	minus	cctCGCCCcgtcg	0.944	0.962	V\$POLR3A_01		>	>		19176527; 9331371				
Factor Name Region Strand Sequence Dissimilarity RE Equally RE Query WT c236 C.>T c336 Factor (C->T Reference (PMID) TFIH 664669 plus TTCTCC 11.337888 1.24512 1.92792 C90 G.>C C.>T Reference (PMID) TFIH 664669 plus TTCTCC 11.337888 1.24512 1.9483 C90 G.>C C.>T ' 17239664 PXX-5 805811 plus GCCGCCCC 9.552105 0.74707 1.9483 ' ' ' ' ' 22278416 TP33 805811 plus GCCGCCCC 9.552105 0.74707 1.9483 ' ' ' ' ' ' 22278416 ' <</td <td></td> <td></td> <td></td> <td></td> <td></td> <td>Allgene</td> <td>PROMO</td> <td></td> <td></td> <td></td> <td></td> <td></td>						Allgene	PROMO									
MatchFactor NameRegionStrandSequenceDissimilarityRE FquallyRE QueryWT $C250$ $C50$ $C50$ C-260TFIH 664669 plusTTCTCC 11.337888 1.24512 1.92792 $C90$ $C250$ $C.71$ Reference (PMID)TFIH 664669 plusTTCTCC 11.337888 1.24512 1.92792 $C90$ $C.2378416$ PXX5 805811 plusGCCGCCC 9.552105 0.74707 1.9483 $C250$ $C.71$ $C.2278416$ TP53 805811 plusGCCGCCCC 9.552105 0.74707 1.9483 $C.77$ $C.2278416$ SP1 803812 plusGCGCGCCCC 3.170707 0.01751 0.09628 V V $C.26439195;$ SP1 811815 plusCCTCG 8.281568 3.98438 5.91804 V V V $26439195;$ SP1 811815 plusCCTCG 8.281568 3.98438 5.91804 V V V $Z0778979;$ SP1 811815 plusCCTCG 8.281568 3.98438 5.91804 V V V $Z0778979;$									E C C C C	00)					
TFIH 664.669 plus TTCTCC 11.37888 1.24512 1.92792 / <th <="" th=""> / <th <="" th=""> <th <="" th=""> <th <="" th=""></th></th></th></th>	/ / <th <="" th=""> <th <="" th=""> <th <="" th=""></th></th></th>	<th <="" th=""> <th <="" th=""></th></th>	<th <="" th=""></th>		Factor Name	Region	Strand	Sequence	Dissimilarity	RE Equally	RE Ouerv	WT	c200 C->1 c90 G->C	G-30	C:-200 C>T	Reference (PMID)
PAX5 805.811 plus GCCGCCC 9.552105 0.74707 1.9483 7 7 2.2278416 TP53 805.811 plus GCCGCCC 9.552105 0.74707 1.9483 7 7 7 7 7 7 7 22278416 TP53 805.811 plus GCCGCCC 9.552105 0.74707 1.9483 7 7 26439195; 25407019 SP1 803.812 plus GCGCGCCCC 3.170707 0.01751 0.09628 7 7 26078979; 25352121 GRalpha 811815 plus CCTCG 8.281568 3.98438 5.91804 7 7 22078416	TFILI	664 669	sult	LTCTCC	11 337888	1 24512	1 97797		1			17230664				
TP53 805811 plus GCCGCCC 6.188498 0.31128 0.91106 / / 26439195 25407019 SP1 803812 plus CGGCGCCC 3.170707 0.01751 0.09628 / / / 26439195 25352121 SP1 803812 plus CGGCGCCCC 3.170707 0.01751 0.09628 / / / 26078979 25352121 GRalpha 811815 plus CCTCG 8.281568 3.98438 5.91804 / / 23211818	PAX-5	805811	plus	GCCGCCC	9.552105	0.74707	1.9483		. `	`		22278416				
SP1 803812 plus CGGCCGCCCC 3.170707 0.01751 0.09628 v v 26078979 25352121 GRalpha 811815 plus CCTCG 8.281568 3.98438 5.91804 v v v 23211818	TP53	805811	plus	GCCGCCC	6.188498	0.31128	0.91106		>	>		26439195; 25407019				
GR-alpha 811815 plus CCTCG 8.281568 3.98438 5.91804 / / 23211818	SP1	803812	plus	CGGCCGCCCC	3.170707	0.01751	0.09628		`	>		26078979; 25352121				
	GR-alpha	811815	plus	CCTCG	8.281568	3.98438	5.91804		>	>		23211818				
	OI UNC INAULA. W	nule une lowe	T Case Jerier	S FEFET LO DOSIUORS LITAL IL	DATCH to the remain	ID2 DAFL OF LICE II	ALTIX: COTC SUMMARY	V. LIEC CU	The SHITTLE SCORE	C TOT THE	JALLIX IIIAU	CD (The maurix core is denned as the				

for consecutive most conserved nucleotides within the matrix); Matrix similarity the matrix similarity score for the matrix match. The Match score can vary from 0 (lowest similarity) to 1 (highest similarity) of the matrix consecutive most conserved nucleotides within the matrix; match to result, for which the core and matrix similarity are higher than the chosen cutoffs; Matrix ID, identifier for the matrix with considered: (1) Equiprobability for the 4 nucleotides (RE equally), and (2) Estimate the nucleotide probability as the nucleotide frequencies in the query sequence (RE query). Results from these two software analyses highlight the appearance or disappearance of specific TFs binding sites in the presence of no analyzed mutations (wild-type column), both examined ones (c.-236 C>T, c.-90 G>C column), and single separated variants (respectively, c.-90 G>C column). which the putative binding site was found; Dissimilarity the dissimilarity threshold controls how similar a sequence must be to the matrix to be reported as a hit. It was set at 15% (85% similarity. Random expectation (RE), gives the number of expected occurrences of the match in a random sequence of the same length as the query sequence according to the dissimilarity index. Two models are

Two Promoter Variants Associated With Stargardt Phenotype

Investigative Ophthalmology & Visual Science

Two Promoter Variants Associated With Stargardt Phenotype

FIGURE 5. Cytoscape images from network views. This shows the Cytoscape pathways analysis, supported by GeneMANIA and MCODE plug-ins, with nodes and edges reflecting relationships between query TFs involved in *ELOVL4* expression. Edge colors: Coexpression (*light purple*), physical interaction (*antique pink*), genetic interaction (*green*), shared protein domains (*golden yellow*), pathway (*light blue*), predicted (*orange*), and common function (*gray*). (**A**) Graphic representation of all TFs in exam relationship. (**B-E**). MCODE clustering analysis, which evidences that most of the query TFs could be grouped into four functional and/or structural clusters.

alpha^{50,51}) and repressors (ZF5, ⁵² EZF-6, ⁵³ CPBP, ⁵⁴⁻⁵⁶ BCL6B, ⁵⁷ GKLF, ⁵⁸⁻⁶⁰ and RXR-alpha⁶¹⁻⁶⁴).

We can speculate that, as emerges from Cytoscape pathway analysis, the *ELOVL4* expression reduction in the proband could derive from a complex balance of all TFs, most of which could present a mutual influence in determining the final effect. Focusing on TFs binding sites formed as a result of the presence of both examined variants, SP1 probably represents

Science	
& Visual &	
Dphthalmology 8	
Investigative C	

Ē
g
Ę
5
Ę
ğ
5
P
4
5
Q
EI
5
II
Þ
- N
ō
2
1.5
Ë
Ę
0
Sis
Ą
na
4
av
ß
II.
ď
Š
at
š
ž
0
2
3LE
3

Gene NameMonotifies (CD)Proti IDConstrained (CD)Proti IDConstrained (CD)Note IDNote ID <th< th=""><th></th><th></th><th>Ensembl</th><th>Entrez</th><th>RefSeq</th><th></th><th></th><th>Log</th><th>MCODE</th><th>MCODE</th><th>MCODE</th></th<>			Ensembl	Entrez	RefSeq			Log	MCODE	MCODE	MCODE
U1COORDESC CONDUCTON CONDUCTONENEROTON CONDUCTON CO	Gene Name	Annotations (GO ID)	Protein ID	Gene ID	mRNA ID	UniProt ID	Score	Score	Node_Status	Cluster	Score
Life Control C	KLF4	GO:0048598[GO:0001159]GO:000076[GO:0000790] GO:0019827[GO:0044212[GO:0000981]GO:0000975 GO:0000785[GO:0043565]GO:0001085[GO:0044454] GO:0000987[GO:000228]GO:0001228]	ENSP0000404922	9314	NM_004235	043474	0.653	-0.426	Clustered	Cluster 4	3.692
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	KLF6	GO:0003690 GO:0003700 GO:0005515 GO:0046872	ENSP00000445301	1316	NM_001300	Q99612	0.670	-0.400	Clustered	Cluster 3	5.333
$ \begin{array}{c} \ \mbox{Cl} $	E2F6	GO:0003677 GO:0003700 GO:0003714 GO:0005515	ENSP00000446315	1876	NM_212540	075461	0.605	-0.502	Seed	Cluster 4	3.733
IIIConsolies/ICo	GTF21	GO:0006367 GO:0006352	ENSP00000460070	2969	NM_033001	P78347	0.562	-0.577	Clustered	Cluster 2	4.200
RXR COMMANT-ISCOMMANT-RECO	TAFI	GO:0006367[GO:0000975[GO:0003713][GO:0043565] GO:0006352[GO:0016570]GO:0044212[GO:0002039] GO:0011067	ENSP00000424526	6872	NM_138923	TAF1_HUMAN	0.719	-0.330	Clustered	Cluster 2	4.167
AndConsist of consist (COMO)Consist (COMO)Consi	1000	CO.0000011CO.00063671CO.00000751CO.000072851	CLOOP/00000000	7267	720000 MIN	DVDA TITIMAN	2120	1010	Chrotomod	Custon	000 %
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	KAKA	GO:00003713 GO:000790 GO:0006352 GO:0044454 GO:00003713 GO:000790 GO:0006352 GO:0044454 GO:0000228 GO:0044212 GO:0001067	ENST000004/0812	0670	/ 66700 TWN	KAKA_HUMAN	010.0	-0.484	Clustered	Cluster 2	4.000
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	ZBTB7A	GO:0000978 GO:0001078 GO:0003677 GO:0003700 GO:0005515	ENSP00000471865	51341	NM_015898	ZBT7A_HUMAN	0.770	-0.262	Unclustered	~	2.311
$M35$ $CO000078]CO000077](CO000570]$ ENEPD00047057 $S07$ M_0 0.574 0.354 0.702 0.354 0.644 <th< td=""><td>HDAC2</td><td>GO:0048598[GO:0000785]GO:0043565[GO:0001085] GO:0000790[GO:0044454[GO:0000228[GO:0016270</td><td>ENSP00000430432</td><td>3066</td><td>NM_001527</td><td>Q92769</td><td>0.588</td><td>-0.531</td><td>Clustered</td><td>Cluster 2</td><td>3.889</td></th<>	HDAC2	GO:0048598[GO:0000785]GO:0043565[GO:0001085] GO:0000790[GO:0044454[GO:0000228[GO:0016270	ENSP00000430432	3066	NM_001527	Q92769	0.588	-0.531	Clustered	Cluster 2	3.889
NRS(1 CO000657(0)C000652 ENSPR000472672 2008 NU_00124/56 PH190 0.581 -0.530 Duckneed Clustered Clu	PAX5	GO:000078[GO:0001077[GO:0003677]GO:0003700] GO:0005515	ENSP00000431038	5079	NM_016734	Q02548	0.702	-0.354	Clustered	Cluster 1	8.000
SPIB CO.00075/IGCO.0014312(CO.00035) ESSP000047365.5 ESSP00004756.5 ESSP00004756.5<	NR3C1	GO:0006367 GO:0006352	ENSP00000427672	2908	NM_001204264	P04150	0.583	-0.539	Clustered	Cluster 4	3.714
TP33 GO0000591GC>0000375[GO000158] ENSP00000756[GO000158] ENSP00000756[GO000158] ENSP00000756[GO000158] ENSP00001756[GO000756] ENSP00001756[GO000756] ENSP0000157 GO0000575[GO000128](GO000756] ENSP0000157 GO000575[GO000158](GO000756] ENSP0000458133 6667 NNL J38473 SPL JHUMAN 0.547 -0.603 Clastered	SPIB	GO:0003700 GO:0043565	ENSP00000472626	6689	NM_003121	SPIB_HUMAN	0.644	-0.440	Unclustered		2.700
Groundist Groundist <t< td=""><td>TP53</td><td>GO:0000790 GO:2000377 GO:0044212 GO:0002039 GO:0000981 GO:0000975 GO:000785 GO:0001085 </td><td>ENSP00000473895</td><td>7157</td><td>NM_001276761</td><td>P53_HUMAN</td><td>0.547</td><td>-0.603</td><td>Clustered</td><td>Cluster 2</td><td>4.209</td></t<>	TP53	GO:0000790 GO:2000377 GO:0044212 GO:0002039 GO:0000981 GO:0000975 GO:000785 GO:0001085	ENSP00000473895	7157	NM_001276761	P53_HUMAN	0.547	-0.603	Clustered	Cluster 2	4.209
$SP1$ G0:0006367[G0:0001159[G0:0000575[G0:000057] Exsponde358133 6667 NM_138473 SP1_HUMAN 0.586 -0.534 Seed Currer 2 C0:00043565[G0:0001057][G0:000057] Exsponde473846 5478 NM_006193 PXX4_HUMAN 0.769 -0.262 Clustered // PLX4 G0:000957[G0:000377][G0:00035716] Exsp000045355 7541 NM_00519 20769 -0.315 Unclustered // ZBTB14 G0:000457[G0:000376][G0:0003706][G0:0003706][G0:0003513] Exsp000045356 7020 NM_003220 P05549 0.779 -0.315 Unclustered // ZBTB14 G0:0004356[G0:000376][G0:000376][G0:00037313[C0:0004353] Exsp000045350 7020 NM_003220 P03549 0.779 -0.315 Unclustered // ZBTB12 G0:0004356[G0:000357][G0:0003582] Exsp0000473380 11128 NM_00755 P0.321 Clustered Clustered Clustered // ZBTA2 G0:0004356[G0:000356][G0:000356][G0:0003582][G0:0003582][G0:0003582][G0:000358] Exsp0000473397 RtD NM_00755 RtCL_HUMAN 0.807		GO:0044454 GO:0000228 GO:0001228 GO:0016570 GO:0001067									
$ \begin{array}{c} C0.004356 GO.000635 GO.000637 GO.000637 GO.000637 GO.000636 GO.000637 GO.0004358 GO.0004358 GO.0004356 GO.0004358 GO.0004356 GO.00037 GO.00037 GO.0004358 GO.0004358 GO.0004358 GO.0004358 GO.0004358 GO.0004358 GO.00037 GO.00037 GO.00037 GO.00037 GO.00037 GO.00037 GO.00037 GO.0004358 GO.0004358 GO.0004358 GO.00037 GO.0003$	IdS	GO:0006367 GO:0001159 GO:0000975 GO:0000976	ENSP00000458133	6667	NM_138473	SP1_HUMAN	0.586	-0.534	Seed	Cluster 2	4.678
P1X4 GO:000980[GO:0001206](GO:0003677][GO:000350] ENSP00000473846 5078 NM_00519 DXX_IHUMAN 0.769 -0.262 Clustered Clustered / ZB7B14 GO:000977][GO:00003716](GO:0005515] ENSP0000463555 7541 NM_003409 ZB714_HUMAN 0.730 -0.315 Unclustered / ZB7B14 GO:00043563 7521 ENSP0000465355 7541 NM_003200 ZB714_HUMAN 0.730 -0.315 Unclustered / ZB724 GO:0043563 GO:0043563 7020 NM_003220 P05549 0.725 -0.321 Clustered Clustered ZB476 GO:0043565 GO:0043576 ENSP000042358 7020 NM_00755 RPC_I_HUMAN 0.730 -0.214 Unclustered / GO:0004370 GO:000357[GO:0003582[GO:0003682[GO:0003682]GO:0003682[GO:000473389 11128 NM_007055 RPC_I_HUMAN 0.817 -0.214 Unclustered / GO:0004370 GO:000367[GO:0003682[GO:0003682[GO:0003682]GO:0003682[GO:0003682]GO:0003682]GO:000367[GO:0003682]GO:0003682]GO:0003672]BO SPG PO PO </td <td></td> <td>GO:0043565 GO:0001085 GO:0006352 GO:000987 GO:0042826 GO:0044212 GO:0001067</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>		GO:0043565 GO:0001085 GO:0006352 GO:000987 GO:0042826 GO:0044212 GO:0001067									
ZBTB14 GC:000097](GC:0000981]GC:0005515] ENSP00000463555 7541 NM_003409 ZBT14_HUMAN 0.730 -0.315 Unclustered $/$ TFAP2A GC:00043576](GC:000097](GC:0004321] ENSP00000420568 7020 NM_003220 $P05549$ 0.725 -0.321 Clustered $/$ GC:0004375[GC:000097](GC:0004375](GC:0004328] ENSP00004205 7020 NM_003220 $P05549$ 0.725 -0.321 Clustered $/$ GC:0004375[GC:000377](GC:0004375] ENSP000047539 11128 NM_007055 RPC1_HUMAN 0.807 -0.214 Unclustered $/$ $POLR34$ GC:0000377[GC:000367](GC:000357] ENSP00000472397 8463 NM_007055 RPC1_HUMAN 0.807 -0.214 Unclustered $/$ $POLR34$ GC:00003570 ENSP00000472397 8463 NM_007055 RPC1_HUMAN 0.807 -0.214 Unclustered $/$ $POLR34$ GC:00003570 ENSP00000472397 8463 NM_003598 TEAD2_HUMAN 0.807 -0.214 Unclustered <t< td=""><td>PAX4</td><td>GO:0000980 GO:0001206 GO:0003677 GO:0003690 GO:0005515</td><td>ENSP00000473846</td><td>5078</td><td>NM_006193</td><td>PAX4_HUMAN</td><td>0.769</td><td>-0.262</td><td>Clustered</td><td>Cluster 1</td><td>8.000</td></t<>	PAX4	GO:0000980 GO:0001206 GO:0003677 GO:0003690 GO:0005515	ENSP00000473846	5078	NM_006193	PAX4_HUMAN	0.769	-0.262	Clustered	Cluster 1	8.000
$ TE4P2A = G0.0045598[G0.0001159[G0.000976]G0.001531] = NSP00000420568 7020 NM_003220 P05549 0.725 -0.321 Clustered Cluster 2 G0.0042472[G0.004839]G0.2000987[G0.0044212] G0.0044212] G0.0042471[G0.004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.0004356]G0.000367[G0.0004329] = NNP00000473389 11128 NM_007055 RPC1_HUMAN 0.807 -0.214 Undustered / G0.0001056]G0.000357[G0.000367]G0.000362] = NSP00000473389 11128 NM_007055 RPC1_HUMAN 0.807 -0.214 Undustered / G0.0000357[G0.000357]G0.000357]G0.000352 = NNP00000472397 8463 NM_00755 RPC1_HUMAN 0.817 -0.203 Undustered / G0.0000357[G0.0006357]G0.0006357[G0.000352] = NNP00000472397 8463 NM_007558 RPC1_HUMAN 0.817 -0.203 Undustered / G0.0000357[G0.0006357]G0.0006357[G0.000472397 8463 NM_007558 RPC1_HUMAN 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.000535[G0.000472397 8463 NM_007558 RPC1_HUMAN 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006357[G0.0006357]G0.0006472397 8463 NM_0007588 TFAD2_HUMAN 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006357[G0.0006357]G0.0006472397 RFE2D2 MANNA 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.000535[G0.0006472397 8463 NM_0075886] TFAD2_HUMAN 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006357[G0.0006472397 TFE2D2 MANNA 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006357[G0.0006472397 TFE2D2 MANNA 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006870 TFE2D2 MANA 0.817 -0.203 Undustered / G0.00008370 TFE2D2 G0.0006870 TFE2D2 MANA 0.817 -0.203 Undustered / G0.00008370 TFE2D2 TFE2D2 MED2 TFE2D2 MANA 0.817 -0.203 Undustered / G0.0000870 TFE2D2 MED2 MED2 MED2 MED2 MED2 MED2 MED2 $	ZBTB14	GO:0000977 GO:0000981 GO:0003700 GO:0005515 GO:0043565	ENSP00000463555	7541	NM_003409	ZBT14_HUMAN	0.730	-0.315	Unclustered	~	3.467
G0:0001067 G0:0003677 [G0:0003682 [G0:0003682 [G0:0003682 [G0:0003682 [G0:0003677] [G0:0003667] [G0:0003667] [G0:0003667] [G0:0003667] [G0:0003682 [G0:0003682 [G0:0003473389] ENSP00000473389 11128 NM_007055 RPCL_HUMAN 0.807 -0.214 Unclustered / G0:0001056 [G0:0003667] [G0:0003682 [G0:0003682 [G0:0003472397] ENSP00000472397 8463 NM_00755 RPCL_HUMAN 0.817 -0.203 Unclustered / G0:00063667 [G0:0006352 ENSP00000472397 8463 NM_003598 TEAD2_HUMAN 0.817 -0.203 Unclustered / Gene name, name of the gene that encodes for one of involved transcription factor. Annotation (G0 ID), ID associated to gene ontology. The GO classifies gene functions along three as molecular function, (2) cellular component, and (3) biological process; Ensembl protein ID, protein ID for Entrez gene ID, gene ID for Entrez; RefSeq mRNA ID, mRNA ID for NCBI Sequence Database; UniProt ID, UniProt protein ID score. Prior to construction, the selected networks are each assigned a weight by the GeneMANIA algorithm. The query genes are assign value of 1, while all other genes are 0. Label propagation is then applied to the entire network and the resulting labels are saved as the score attribute, used to rank the genes. The score attribute to the network and the resulting labels are saved as the score attribute, used to rank the genes. The score decore decore assign form on the network and how long and heavily weighted those path are relevance of each go decore duct and the network and how long and heavily weighted those path are relevare assign form a	TFAP2A	GO:0048598[GO:0001159]GO:0000976[GO:0016331] GO:0042472]GO:0048839[GO:2000377]GO:0044212] GO:000981]GO:000975[GO:0003713]GO:0043583] GO:0043565[GO:000987]GO:0001228[GO:0042471]	ENSP00000420568	7020	NM_003220	P05549	0.725	-0.321	Clustered	Cluster 2	4.182
POLR3A G0:0001056[G0:0003677]G0:0003682[G0:0003899] ENSP00000473399 11128 NM_007055 RPC1_HUMAN 0.807 -0.214 Unclustered / G0:0008270 G0:000352 ENSP0000472397 8463 NM_00755 RPC1_HUMAN 0.807 -0.214 Unclustered / G0:0008270 G0:0006367[G0:0005352 ENSP00000472397 8463 NM_003598 TEAD2_HUMAN 0.817 -0.203 Unclustered / Gene name, name of the gene that encodes for one of involved transcription factor. Annotation (GO ID), ID associated to gene ontology. The GO classifies gene functions along three as molecular function, (2) cellular component, and (3) biological process; Ensembl protein ID, protein ID for Enterz gene ID, gene ID for Enterz; RefSeq mRNA ID, mRNA ID for NCBI Sequence Database; UniProt ID, UniProt protein ID score. Prior to construction, the selected networks are each assigned a weight by the GeneMANIA algorithm. The query genes are assign value of 1, while all other genes are 0. Label propagation is then applied to the entire network and the resulting labels are saved as the score attribute, used to rank the genes. The score assign gene node end up in one of the query nodes and how long and heavily weighted those paths are relevance of each grach decomponent and the resulting labels are saved as the score attribute, used to rank the genes. The score assign gene node end up in one of the query nodes and how long and heavily weighted those paths are relevance of each grach decore assign gene node end up in one of the query nodes		GO:0001067									
TEAD2 GO:0006367 [GO:0006352] ENSP00000472397 8463 NM_003598 TEAD2_HUMAN 0.817 -0.203 Unclustered / Gene name, name of the gene that encodes for one of involved transcription factor. Annotation (GO ID), ID associated to gene ontology. The GO classifies gene functions along three as molecular function, (2) cellular component, and (3) biological process; Ensembli protein ID, protein ID for Ensembli; Entrez gene ID, gene ID for Entrez; RefSeq mRNA ID, mRNA ID for NCBI. Sequence Database; UniProt ID, UniProt protein ID score. Prior to construction, the selected networks are each assigned a weight by the GeneMANIA algorithm. The query genes are assign value of 1, while all other genes are 0. Label propagation is then applied to the entire network and the resulting labels are saved as the score attribute, used to rank the genes. The score atsign to the network and how long and heavily weighted those paths are. This score indicates the relevance of each gradient function to the network cand how long and heavily weighted those paths are. This score indicates the relevance of each gradient function to the network cand how long and heavily weighted those paths are. This score indicates the relevance of each gradient function to the network cand how long and heavily weighted those paths are. This score indicates the relevance of each gradient function to the network cand in the encoder category and the score and the score attribute come control form barbow control of the network cand in the encoder category and the score attribute to the relevance of each gradient function to the network cand in the encoder category attribute to the network category attribute to the network and how long and heavily weighted those paths are relevance of each gradient parts the score attribute to the network category attribute to the network ca	POLR3A	GO:0001056 GO:0003677 GO:0003682 GO:0003899 GO:0008270	ENSP00000473389	11128	NM_007055	RPC1_HUMAN	0.807	-0.214	Unclustered	~	2.000
Gene name, name of the gene that encodes for one of involved transcription factor. Annotation (GO ID), ID associated to gene ontology. The GO classifies gene functions along three as molecular function, (2) cellular component, and (3) biological process; Ensembl protein ID for Ensembl; Entrez gene ID, gene ID for Entrez; Refseq mRNA ID, mRNA ID for NCBI. Sequence Database; UniProt ID, UniProt protein ID score. Prior to construction, the selected networks are each assigned a weight by the GeneMANIA algorithm. The query genes are assign value of 1, while all other genes are 0. Label propagation is then applied to the entire network and the resulting labels are saved as the score attribute, used to rank the genes. The score assign gene reflects how often paths starting at a given gene node end up in one of the query nodes and how long and heavily weighted those paths are. This score indicates the relevance of each g	TEAD2	GO:0006367/GO:0006352	ENSP00000472397	8463	NM_003598	TEAD2_HUMAN	0.817	-0.203	Unclustered	/	1.533
OTPINA ISI DASCU OLI UE SELECICU RELVOTES. FIERICI SCOLES INICALE SCILES UNE REL REL REL REL RUE	Gene nam molecular fun Sequence Data value of 1, whi gene reflects E original list ba	e, name of the gene that encodes for one of involved ction, (2) cellular component, and (3) biological proc abase; UniProt ID, UniProt protein ID score. Prior to ile all other genes are 0. Label propagation is then app tow often paths starting at a given gene node end up sed on the selected networks. Hisher scores indicate	transcription factor. ess; Ensembl protein construction, the sel lied to the entire net n one of the query n	Annotation (ID, protein] ected networ work and the odes and hove	(GO ID), ID assoc ID for Ensembl; Er eks are each assign e resulting labels a w long and heavily functionally relativ	ziated to gene onto. Intrez gene ID, gene ned a weight by the ure saved as the scon y weighted those pi y we also chose pi red. We also chose	logy. The C ID for Ent c GeneMAl c attribute aths are. T not to add	O classific rez; RefSec VIA algorit , used to r nis score ir anv other	s gene functions (mRNA ID, mRN hm. The query g ank the genes. T dicates the relev- tents coming fi	a along three a A ID for NCB cenes are assig he score assign ance of each rom pathway	ispects: (1) Reference ned a label ned to each gene to the analys is to

Two Promoter Variants Associated With Stargardt Phenotype

Error Bars: +/- 1 SD

FIGURE 6. Luciferase assay results. The histogram shows the means coming from luciferase ratios between Firefly and Renilla bioluminescence measurements (in relative luminescence units - LRU) for each sample. As reported, ANOVA test resulted significant ($P = 1.1391^{E-109}$), also for multiple comparisons (highlighted with pairwise lines on *bars* and *asterisk*). The presence of only the c.-90 G>C or the c. 236 C>T variant determines an *ELOVL4* expression reduction, respectively, of approximately 14% or 18%, compared with the healthy control, while the heterozygous condition for both examined variants drastically lowers it (~97%).

the key node around which other factors determine the overall *ELOVL4* protein under expression. As literature evidences, SP1 should be enhanced by RXR-alpha activity,⁶⁵ but this positive status should be repressed or seized by interaction with TFII-I,⁶⁶ TP53,⁶⁷ and KLF6,⁶⁸ the latter stimulated by KLF4.⁶⁹ The direct consequences of SP1 inhibition could be related to a decrease of connected PAX-5,^{70,71} in turn influencing PAX-4.⁷² Although there is no solid proof, in literature, of a possible direct interaction, we can hypothesize, basing on previous assumptions, that TFII-I, usually acting as a repressor, could contribute to the inhibition of SP1, as well as the possibility that SP1 impairment could reflect on POLR3A, reducing its activity.

Due to the unavailability of data on involved TF interactions, analysis on TF binding sites lost in the patient is not clear. We can only speculate that, because HDAC2 usually acts as a repressor, with FBI-I as corepressor,⁷³ both could downregulate another two inhibitors, ZF5 and E2F-6. This situation leaves most of the transcriptional activity to TAFII250, which functions as an activator, and which presents many binding sites in wild-type genotype.

CONCLUSIONS

A reduction of Elovl4 enzymatic function in the endoplasmic reticulum (ER) could result in a deficiency of VLC-PUFA,

which may be required for the construction, function, and maintenance of healthy OS or other photoreceptor membranes; hence, the absence of sufficient quantities eventually results in retinal degeneration. The results presented here reported reduced Elovl4 enzyme activity, fundamental in VLC- PUFA synthesis, vital for rod function and rod longevity, parameters that are involved in the etiopathogenesis of STGD.

We speculate that an altered balance of TF binding sites, due to the presence of c.-90 G>C and c.-236 C>T, and the possible interaction of involved transcription factors, could determine an overall prevalence of repressive activity rather than enhancing activity, resulting in a downregulation of *ELOVL4* expression, as functionally demonstrated by the dualluciferase reporter assay.

Even if the in vitro experiments demonstrate an expression reduction of *ELOVL4* promoter due to transconfiguration of analyzed variants, we cannot assert with certainty that the same effect, in vivo, is limited to both variants' presence. For example, cells are treated outside their normal 'microenvironment' (no surrounding tissues, no blood supply, no normal supply of nutrients, etc.), and we cannot exclude the involvement of other factors into the altered expression of *ELOVL4*. Moreover, further experiments (e.g., ChIP-Sequencing of involved TFs) will be needed to confirm, or not, the role of single transcription factors and reciprocal interactions, involved in *ELOVL4* downregulation.

Acknowledgments

The authors thank all of the study participants.

Disclosure: L. Donato, None; C. Scimone, None; C. Rinaldi, None; P. Aragona, None; S. Briuglia, None; A. D'Ascola, None; R. D'Angelo, None; A. Sidoti, None

References

Investigative Ophthalmology & Visual Science

- 1. Jiang, F, Pan, Z, Xu K, et al. Screening of ABCA4 gene in a chinese cohort with Stargardt disease or cone-rod dystrophy with a report on 85 novel mutations. *Invest Ophthalmol Vis Sci.* 2016;57:145–152.
- Lansel N, Niemeyer G, Thölen A. Stargardt's disease and its intrafamilial variability [in German]. *Klin Monbl Augenbeilkd*. 2000;216:342-345.
- Tanna P, Strauss RW, Fujinami K, Michaelides M. Stargardt disease: clinical features, molecular genetics, animal models and therapeutic options. *Br J Ophthalmol.* 2017;101:25–30.
- Yi J, Li S, Jia X, et al. Evaluation of the ELOVL4, PRPH2 and ABCA4 genes in patients with Stargardt macular degeneration. *Mol Med Rep.* 2012;6:1045-1049.
- Agbaga MP, Tam BM, Wong JS, Yang LL, Anderson RE, Moritz OL. Mutant ELOVL4 that causes autosomal dominant Stargardt-3 macular dystrophy is misrouted to rod outer segment disks. *Invest Ophthalmol Vis Sci.* 2014;55:3669–3680.
- Logan S, Anderson RE. Dominant Stargardt macular dystrophy (STGD3) and ELOVL4. Adv Exp Med Biol. 2014;801:447-453.
- Bardak H, Gunay M, Erçalık Y, et al. Analysis of ELOVL4 and PRPH2 genes in Turkish Stargardt disease patients. *Genet Mol Res.* 2016;15:gmr15048774.
- Tran HV, Moret E, Vaclavik V, et al. Swiss family with dominant Stargardt disease caused by a recurrent mutation in the ELOVL4 gene. *Klin Monbl Augenheilkd*. 2016;233:475-477.
- Harkewicz R, Du H, Tong Z, et al. Essential role of elovl4 protein in very long chain fatty acid synthesis and retinal function. *J Biol Chem.* 2012;287;11469–11480.

- Bennett LD, Anderson RE. Current progress in deciphering importance of VLC-PUFA in the retina. *Adv Exp Med Biol.* 2016;854;145-151.
- 11. Marchette LD, Sherry DM, Brush RS, et al. Very long chain polyunsaturated fatty acids and rod cell structure and function. *Adv Exp Med Biol.* 2014;801:637-645.
- 12. Logan S, Agbaga MP, Chan MD, et al. Deciphering mutant ELOVL4 activity in autosomal-dominant Stargardt macular dystrophy. *Proc Natl Acad Sci U S A*. 2013;110:5446-5451.
- Bennett LD, Hopiavuori BR, Brush RS, et al. Examination of VLC-PUFA-deficient photoreceptor terminals. *Invest Ophthalmol Vis Sci.* 2014;55;4063-4072.
- Ayyagari R, Zhang K, Hutchinson A, et al. Evaluation of the ELOVL4 gene in patients with age-related macular degeneration. *Ophthalmic Genet*. 2001;22:233–239.
- 15. Aguilà M, Bevilacqua D, McCulley C, et al. Hsp90 inhibition protects against inherited retinal degeneration. *Hum Mol Genet*. 2014;23:2164–2175.
- 16. Matys V, Kel-Margoulis OV, Fricke E, et al. TRANSFAC and its module TRANScompel: transcriptional gene regulation in eukaryotes. *Nucleic Acids Res.* 2006;34:D108-D110.
- Farré D, Roset R, Huerta M, et al. Identification of patterns in biological sequences at the ALGGEN server: PROMO and MALGEN. *Nucleic Acids Res.* 2003;31:3651–3653.
- Scimone C, Bramanti P, Ruggeri A, et al. CCM3/SERPINI1 bidirectional promoter variants in patients with cerebral cavernous malformations: a molecular and functional study. *BMC Med Genet.* 2016;17:74.
- Tzeng DT, Tseng YT, Ung M, Liao IE, Liu CC, Cheng C. DPRP: a database of phenotype-specific regulatory programs derived from transcription factor binding data. *Nucleic Acids Res.* 2014;42:D178-D183.
- Liu X, Yu X, Zack DJ, Zhu H, Qian J. TiGER: a database for tissue-specific gene expression and regulation. *BMC Bioinformatics*. 2008;9:271.
- 21. Rheinbay E, Suvà ML, Gillespie SM, et al. An aberrant transcription factor network essential for Wnt signaling and stem cell maintenance in glioblastoma. *Cell Rep.* 2013;3: 1567-1579.
- 22. Myšičková A, Vingron M. Detection of interacting transcription factors in human tissues using predicted DNA binding affinity. *BMC Genomics*. 2012;13(suppl 1):S2.
- 23. Bennett LD, Brush RS, Chan, M, et al. Effect of reduced retinal VLC-PUFA on rod and cone photoreceptors. *Invest Ophtbalmol Vis Sci.* 2014;55:3150–3157.
- Palejwala NV, Gale MJ, Clark RF, Schlechter C, Weleber RG, Pennesi ME. Insights into autosomal dominant Stargardt-like macular dystrophy through multimodality diagnostic imaging. *Retina*. 2016;36;119–130.
- 25. Vasireddy V, Jablonski MM, Khan NW, et al. ELOVL4 5-bp deletion knock-in mouse model for Stargardt-like macular degeneration demonstrates accumulation of elovl4 and lipofuscin. *Exp Eye Res.* 2009;89:905–912.
- Okuda A, Naganuma T, Ohno Y, et al. Hetero-oligomeric interactions of an elovl4 mutant protein: implications in the molecular mechanism of Stargardt-3 macular dystrophy. *Mol Vis.* 2010;16:438-445.
- 27. Kageyama R, Merlino GT, Pastan I. Nuclear factor ETF specifically stimulates transcription from promoters without a TATA box. *J Biol Chem.* 1989;264;15508–15514.
- Maeda T, Ito K, Merghoub T, et al. LRF is an essential downstream target of GATA1 in erythroid development and regulates BIMdependent apoptosis. *Dev Cell*. 2009;17;527–540.

- 29. Fan J, Alsarraf O, Dahrouj M, et al. linhibition of HDAC2 protects the retina from ischemic injury. *Invest Ophthalmol Vis Sci.* 2013;54:4072-4080.
- Kong X, Fang M, Li P, Fang F, Xu Y. HDAC2 deacetylates class II transactivator and suppresses its activity in macrophages and smooth muscle cells. *J Mol Cell Cardiol.* 2009;46:292–299.
- 31. Koriyama Y, Takagi Y, Chiba, K, et al. Requirement of retinoic acid receptor β for genipin derivative-induced optic nerve regeneration in adult rat retina. *PLoS One*. 2013;8:e71252.
- 32. Lebrun-Julien F, Suter U. Combined HDAC1 and HDAC2 depletion promotes retinal ganglion cell survival after injury through reduction of p53 target gene expression. *ASN Neuro*. 2015;7:pii:1759091415593066.
- Herzfeld T, Nolte D, Müller U. Structural and functional analysis of the human TAF1/DYT3 multiple transcript system. *Mamm Genome*. 2007;18:787-795.
- 34. Piper M, Dwivwdy A, Leung L, Bradley RS, Holt CE. NFprotocadherin and TAF1 regulate retinal axon initiation and elongation in vivo. *J Neurosci*. 2008;28:100–105.
- 35. Ray D, Bosselut R, Ghysdael J, Mattei MG, Tavitian A, Moreau-Gachelin F. Characterization of Spi-B, a transcription factor related to the putative oncoprotein Spi-1/PU.1. *Mol Cell Biol.* 1992;12:4297–4304.
- 36. Rath MF, Bailey MJ, Kim JS, Coon SL, Klein DC, Møller M. Developmental and daily expression of the PAX4 and PAX6 homeobox genes in the rat retina: localization of PAX4 in photoreceptor cells. *J Neurochem.* 2009;108:285–294.
- Reichman S, Kalathur RK, Lambard S, et al. The homeobox gene CHX10/VSX2 regulates RDCVF promoter activity in the inner retina. *Hum Mol Genet*. 2010;19:250–261.
- 38. Natalizio BJ, Robson-Dixon ND, Garcia-Blanco MA. The carboxyl-terminal domain of RNA polymerase II is not sufficient to enhance the efficiency of pre-mRNA capping or splicing in the context of a different polymerase. *J Biol Chem*. 2009;284:8692–8702.
- 39. Sepehri S, Hernandez N. The largest subunit of human RNA polymerase III is closely related to the largest subunit of yeast and trypanosome RNA polymerase III. *Genome Res.* 1997;7: 1006–1019.
- Grueneberg DA, Henry RW, Brauer A, et al. A multifunctional DNA-binding protein that promotes the formation of serum response factor/homeodomain complexes: identity to TFII-I. *Genes Dev.* 1997;11:2482-2493.
- Palmer SJ, Tay ES, Santucci N, et al. Expression of Gtf2ird1, the Williams syndrome-associated gene, during mouse development. *Gene Expr Patterns*. 2007;7:396–404.
- Bougel S, Renaud S, Braunschweig R, et al. PAX5 activates the transcription of the human telomerase reverse transcriptase gene in B cells. *J Pathol.* 2010;220:87–96.
- 43. Livide G, Epistolato MC, Amenduni M, et al. Epigenetic and copy number variation analysis in retinoblastoma by MS-MLPA. *Pathol Oncol Res.* 2012;18:703-712.
- 44. Polato F, Rusconi P, Zangrossi S, et al. DRAGO (KIAA0247), a new DNA damage-responsive, p53-inducible gene that cooperates with p53 as oncosuppressor. [Corrected]. *J Natl Cancer Inst.* 2014;106:dju053.
- 45. Wang Y, Wong MM, Zhang X, Chiu SK. Ectopic AP4 expression induces cellular senescence via activation of p53 in long-term confluent retinal pigment epithelial cells. *Exp Cell Res.* 2015;339:135-146.
- 46. Yasuda M, Tanaka Y, Nishiguchi KM, et al. Retinal transcriptome profiling at transcription start sites: A cap analysis of gene expression early after axonal injury. *BMC Genomics*. 2014;15:982.

- 47. Alfonso-Jaume MA, Mahimkar R, Lovett DH. Co-operative interactions between NFAT (nuclear factor of activated t cells) c1 and the zinc finger transcription factors Sp1/Sp3 and Egr-1 regulate MT1-MMP (membrane type 1 matrix metalloproteinase) transcription by glomerular mesangial cells. *Biochem J*. 2004;380:735-747.
- 48. Bikbova G, Oshitari T, Baba T, Yamamoto S. Altered expression of NF- κB and Sp1 after exposure to advanced glycation end-products and effects of neurotrophic factors in ages exposed rat retinas. *J Diabetes Res.* 2015;2015: 543818.
- 49. Donovan K, Alekseev O, Qi X, Cho W, Azizkhan-Clifford J. O-GlcNAc modification of transcription factor Sp1 mediates hyperglycemia-induced VEGF-A upregulation in retinal cells. *Invest Ophthalmol Vis Sci.* 2014;55:7862-7873.
- Cubilla MA, Bermùdez V, Marquioni Ramella MD, Bachor TP, Suburo AM. Mifepristone, a blocker of glucocorticoid receptors, promotes photoreceptor death. *Invest Ophthalmol Vis Sci.* 2013;54:313-322.
- 51. Nader N, Chrousos GP, Kino T. Circadian rhythm transcription factor clock regulates the transcriptional activity of the glucocorticoid receptor by acetylating its hinge region lysine cluster: potential physiological implications. *FASEB J.* 2009; 23:1572–1583.
- 52. Orlov SV, Kuteykin-Teplyakov KB, Ignatovich IA, et al. Novel repressor of the human FMR1 gene identification of p56 human (GCC)(n)-binding protein as a krüppel-like transcription factor zf5. *FEBS J.* 2007;274:4848–4862.
- 53. Trimarchi JM, Firechild B, Verona R, Moberg K, Andon N, Lees JA. E2F-6, a member of the E2F family that can behave as a transcriptional repressor. *Proc Natl Acad Sci U S A*. 1998;95: 2850–2855.
- 54. Koritschoner NP, Bocco JL, Panzetta-Dutari GM, Dumur CI, Flury A, Patrito LC. A novel human zinc finger protein that interacts with the core promoter element of a TATA box-less gene. J Biol Chem. 1997;272:9573–9580.
- 55. Nakamura H, Edward DP, Sugar J, Yue BY. Expression of Sp1 and KLF6 in the developing human cornea. *Mol Vis.* 2007;13: 1451-1457.
- 56. Steketee MB, Oboudiyat C, Daneman R, et al. Regulation of intrinsic axon growth ability at retinal ganglion cell growth cones. *Invest Ophthalmol Vis Sci.* 2014;55;4369-4377.
- Lim JQ, Lu J, He BP. Diva/BclB regulates differentiation by inhibiting NDPKB/Nm23H2-mediated neuronal differentiation in pc-12 cells. *BMC Neurosci*. 2012;13:123.
- Fang J, Shaw PX, Wang Y, Goldberg JL. Krüppel-Like Factor 4 (KLF4) is not required for retinal cell differentiation. *eNeuro*. 2016;3;pii:ENEURO.0117-0115.2016.
- 59. Rowland BD, Peeper DS. KLF4, p21 and context-dependent opposing forces in cancer. *Nat Rev Cancer*. 2006;6:11-23.
- 60. Villarreal G Jr, Zhang Y, Larman HB, Gracia-Sancho J, Koo A, García-Cardeña G. Defining the regulation of KLF4 expression and its downstream transcriptional targets in vascular endothelial cells. *Biochem Biophys Res Commun.* 2010; 391:984–989.
- Chen H, Privalsky ML. Cooperative formation of high-order oligomers by retinoid x receptors: an unexpected mode of DNA recognition. *Proc Natl Acad Sci U S A*. 1995;92:422– 426.
- 62. Chen Y, Wang W, Liu F, Tang L, Tang R, Li W. 9-cis-retinoic acid improves sensitivity to platelet-derived growth factor-BB via RXRα and SHP-1 in diabetic retinopathy. *Biochem Biophys Res Commun.* 2015;465:810–816.
- 63. Cvekl A, Wang WL. Retinoic acid signaling in mammalian eye development. *Exp Eye Res.* 2009;89:280–291.

Investigative Ophthalmology & Visual Science

- 64. Janssen JJ, Kuhlmann ED, van Vugt AH, et al. Retinoic acid receptors and retinoid X receptors in the mature retina: Subtype determination and cellular distribution. *Curr Eye Res.* 1999;19:338-347.
- 65. Suzuki Y, Shimada J, Shudo K, Matsumura M, Crippa MP, Kojima S. Physical interaction between retinoic acid receptor and Sp1: mechanism for induction of urokinase by retinoic acid. *Blood.* 1999;93:4264-4276.
- 66. Bu Y, Gao L, Gelman IH. Role for transcription factor TFII-I in the suppression of SSeCKS/Gravin/Akap12 transcription by Src. *Int J Cancer.* 2011;128:1836-1842.
- Kong X, Peng B, Yang Y, et al. P53 represses transcription of RING finger LIM domain-binding protein RLIM through Sp1. *PLoS One.* 2013;8:e62832.
- Kong LM, Yao L, Lu N, et al. Interaction of KLF6 and Sp1 regulates basigin-2 expression mediated proliferation, invasion and metastasis in hepatocellular carcinoma. *Oncotarget*. 2016;7:27975-27987.

- 69. Okano J, Opitz OG, Nakagawa H, Jenkins TD, Friedman SL, Rustgi AK. The Krüppel-like transcriptional factors Zf9 and GKLF coactivate the human keratin 4 promoter and physically interact. *FEBS Lett.* 2000;473:95-100.
- Miranda GA, Villalvazo M, Galic Z, et al. Combinatorial regulation of the murine RAG-2 promoter by Sp1 and distinct lymphocyte-specific transcription factors. *Mol Immunol.* 2002;38:1151-1159.
- 71. Wu CX, Zhao WP, Kishi H, et al. Activation of mouse RAG-2 promoter by Myc-associated zinc finger protein. *Biochem Biophys Res Commun.* 2004;317:1096-1102.
- Chalepakis G, Gruss P. Identification of DNA recognition sequences for the PAX3 paired domain. *Gene*. 1995;162:267– 270.
- 73. Choi WI, Jeon, BN, Yun CO, et al. Proto-oncogene FBI-1 represses transcription of p21CIP1 by inhibition of transcription activation by p53 and Sp1. *J Biol Chem.* 2009;284: 12633-12644.