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# Identifying mutations in Tunisian families with retinal dystrophy

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Retinal dystrophies (RD) are a rare genetic disorder with high genetic heterogeneity. This study aimed at identifying disease-causing variants in fifteen consanguineous Tunisian families. Full ophthalmic examination was performed. Index patients were subjected to IROme analysis or whole exome sequencing followed by homozygosity mapping. All detected variations were confirmed by direct Sanger sequencing. Mutation analysis in our patients revealed two compound heterozygous mutations p.(R91W);(V172D) in RPE65, and five novel homozygous mutations: p.R765C in CNGB1, p.H337R in PDE6B, splice site variant c.1129-2A > G and c.678\_681delGAAG in FAM161A and c.1133 + 3\_1133 + 6delAAGT in CERKL. The latter mutation impacts pre-mRNA splicing of CERKL. The other changes detected were six previously reported mutations in CNGB3 (p.R203\*), ABCA4 (p.W782\*), NR2E3 (p.R311Q), RPE65 (p.H182Y), PROM1 (c.1354dupT) and EYS (c.5928-2A > G). Segregation analysis in each family showed that all affected individuals were homozygotes and unaffected individuals were either heterozygote carriers or homozygous wild type allele. These results confirm the involvement of a large number of genes in RD in the Tunisian population.

Retinal dystrophies (RD) are a heterogeneous group of diseases in which the photoreceptor and RPE cells of the retina degenerate, leading to partial or complete blindness and affecting approximately 1 in 2500–3500 individuals<sup>1</sup>. Classification of RD is generally based on whether the disease primarily affects cones or rods (predominantly affecting the macula or peripheral retina respectively) and whether it occurs alone (non-syndromic RD) or in conjunction with other systemic disorders, especially loss of hearing (syndromic RD).

RD phenotypes are variable in terms of onset, progression and severity. The disease may be mild and non progressive, such as in congenital stationary night blindness (CSNB), characterized by defective rod photoreceptors involved in night vision. Non progressive disorders may lead to severe visual impairment as well as achromatopsia (ACHM), stationary congenital cone dystrophies. Other disorders are progressive, leading to severe visual impairment such as in retinitis pigmentosa (RP), cone dystrophy (CD), cone-rod dystrophy (CRD)<sup>2</sup> and Stargardt disease (STGD)<sup>3</sup>. In RP, rod photoreceptors are initially affected more severely than cones<sup>4</sup>. The most severe cases are Leber Congenital Amaurosis (LCA) and early-onset rod-cone dystrophies, in which infants suffer from complete blindness from birth or within the first years of life<sup>5</sup>. Cone dystrophies are characterized by progressive degeneration of cone photoreceptors with preservation of rod function<sup>6</sup>, whereas in CRD peripheral vision is also compromised, leading to early blindness. Stargardt disease is a juvenile macular degeneration characterized by central vision loss<sup>3</sup>.

This group of rare genetic disorders shows substantial clinical and genetic overlaps with high genetic heterogeneity involving more than 220 genes identified so far (https://sph.uth.edu/retnet/). Similar phenotypes may result from different gene mutations, and subtle differences in phenotypes may result from a similar mutation<sup>7</sup>.

The mutations in causative genes induce either in degeneration or dysfunction of retinal cells. Functionally, different gene products are involved in many cellular functions and fall into four categories: proteins directly involved in the phototransduction cascade, genes encoding proteins responsible for the structure and polarity of the photoreceptors, genes encoding proteins of the visual cycle, and regulatory genes (such as transcription and splicing factors)<sup>2</sup>.

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All modes of mendelian inheritance have been described in RD, with autosomal recessive being the most prevalent<sup>2</sup>. In our study we focused on the Tunisian population, known to have a relatively high level of consanguineous marriages, leading to a relatively high frequency of autosomal recessive diseases. This study was designed to apply homozygosity mapping in 15 consanguineous Tunisian families segregating retinal degenerative disease and three families analyzed by IROme<sup>7</sup>, aiming at the identification of the genetic defects.

#### **Materials and Methods**

**Subjects.** The department B of Hedi Rais Institute of Ophthalmology in Tunisia recruited all subjects involved in the study over a 10-year period. 177 Tunisian families segregating RD were enrolled. In this pilot study, a subset of 61 individuals (32 affected, 29 unaffected) from 15 families with clear recessive transmission and two or more affected individuals was selected. The affected individuals comprised 17 males and 15 females, ranging in age from 4 to 72 years, with an onset of disease ranging from birth to 47 years. Demographic characteristics, age at onset, and personal and family history were recorded for all participants. Written informed consent was obtained from each study participant, analyses were done in accordance with local guidelines and regulations study was approved by the Local Ethics Committee of the Hedi Rais Institute.

All patients underwent a standard ophthalmological examination including determination of best corrected visual acuity using standard Snellen charts. Clinical examination was supplemented by fundus photography, color vision assessment using the Farnsworth-Munsell 100 hue color vision test (FM100, Munsell Color Company Inc., Baltimore, MD, USA) and analysis of dark adaptation. Goldmann kinetic perimetry (Carl Zeiss Meditec Inc., Dublin, CA, USA) using V-4e and I-4e targets, fluorescein angiography (Imagenet; Topcon Corporation, Tokyo, Japan) and optical coherence tomography (SD-OCT) (OCT 3D TOPCON 2000) was also performed. Electrophysiological investigation was carried out using the Métrovision vision monitor (Métrovision, Pérenchies, France) according to the International Society for Clinical Electrophysiology of Vision (ISCEV) protocol.

**Homozygosity mapping and mutation analysis.** Genomic DNA of participating individuals was isolated from peripheral blood using a standard salting out procedure. One index patient of 12 families was selected for whole exome sequencing (WES) analysis and unaffected family members were collected for co-segregation analysis. Three index patients were analyzed with IROme<sup>8</sup>.

Exome capturing was performed with the Roche Nimble-Gen version 2 (44.1–megabase pair) at Otogenetics Corporation (Norcross, Georgia) and sequencing was done on an Illumina HiSeq2000 at a mean coverage  $\times$  31. Sequence reads were aligned to the human genome reference sequence (build hg19), and variants were identified and annotated using the Nextgene software package v.2.3.5. (Softgenetics, State College, PA). Homozygosity was evaluated from SNPs obtained by WES. In families with CERKL mutations, SNPs: rs4667591, rs1050354, rs4894140, rs1143676, rs3754929, rs1400130, rs12476147, rs17228441, rs1800255, rs1225090 were selected and analyzed by Sanger sequencing in the index patients of families F12-F15.

Assessment of the Pathogenicity of Candidate Variants. We applied various bioinformatic tools to filter variants against NCBI dbSNP to facilitate the identification of disease-causing mutations. The Exome Aggregation Consortium (ExAC; http://exac.broadinstitute.org) data set was used as a reference data set of population-based allele frequencies $^9$ . Novel DNA variations were compared with data from public databases (HGMD $^{10}$  and 1000 genomes $^{11}$ ). Although most causative mutations associated with RD are rare (MAF <0.01). The putative pathogenicity of the novel missense variants reported in this study was evaluated using two in silico tools: Polymorphism Phenotyping v2 (PolyPhen-2) $^{12}$  and Sorting Intolerant from Tolerant (SIFT) $^{13}$ . The PolyPhen-2 score ranges from 0.0 (tolerated) to 1.0 (deleterious). Variants with scores of 0.0 are predicted to be benign. Values of 1.0 or close to 1.0 are more confidently predicted to be deleterious. PolyPhen-2 and SIFT scores use the same range, 0.0 to 1.0, but with opposite meanings. A variant with a SIFT score of 1.0 is predicted to be benign. In addition, we determined with PolyPhen-2 a measure of evolutionary nucleotide conservation.

**Mutation analysis.** After all filtering steps, Sanger sequencing was used to verify each predicted disease-causing variants remaining in the autozygous regions of interest and to perform cosegregation analyses. PCR was realized in a total reaction mixture of 20 µl, containing 20 ng of genomic DNA, 10 pmol of each primer (Eurogentec, Liège, Belgium) and 10 µl of FastStart PCR Master Mix (Roche, Basel, Switzerland). Three amplification steps (30 s each), with the annealing temperature ranging from 58 °C to 60 °C, were performed. Primer sequences were designed using the PRIMER3 software (http://frodo.wi.mit.edu/) and PCR conditions are provided in Table 1. The purified PCR products were then directly sequenced on an ABI 3100XL DNA automated sequencer (Applied Biosystems, Foster city, CA), using the Big Dye Terminator Labeling Kit version 1 (Life Technologies, Zug, Switzerland). Cosegregation analysis was performed in all families.

**Splicing variant analysis.** A minigene analysis was done to predict the putative impact of the deletion c.1133 + 3\_1133 + 6delAAGT occurring in a splice site. The mutated region was amplified by PCR from DNA of patient VI-1, VI-2 in F13 and one healthy control using specific (forward: 5'-TCC AAA GGA TCC GGG AAC TGA AGA GAA AGT GAG G-3' and reverse: 5'-TCC AAA GAA TTC GGC TGA GTG AGT AGT TGT TTG C-3'). The resulting PCR products were subsequently cloned into the pBK-CMV vector using the T4 DNA Ligase according to manufacturer's protocol (Rapid DNA Ligation Kit). Plasmids were analyzed by direct Sanger sequencing and then transfected into HEK293 cells. Total RNA was extracted with TRIzol (Invitrogen product name), retrotranscribed with the AffinityScript RT (AffinityScript QPCR cDNA Synthesis Kit – Agilent), and the resulting cDNA was PCR-amplified using the previously described primers but without the restriction enzyme sites (forward: 5'-GGG AAC TGA AGA GAA AGT GAG G-3') and (reverse: 5'-GGC TGA GTG AGT AGT TGT TTG C-3'). The amplified products were separated by electrophoresis on a 2% agarose gel and were subsequently analyzed by Sanger sequencing.

Genes	Exon	Forward	Reverse	Hybridation temp. (°C)	
CNGB3	5	GTGAGAACATGCGGTGTTTG	CAAAGATGGGCAAATGATCC	58	
CNGB1	23	AGAGACTCCGCCTCTCACTC GGGGCAGACACGAAGATG		58	
ABCA4	15	AGCACATGGAGTGTGCGTAG	TGCCCCTGTACATTTTAGCC	58	
NR2E3	6	TCTGAGCCTCTGGCTGATGTCA	AGAAGGGAGTCCAGCCTCAC	60	
RPE65	4	GACTTGATGAGGACACATAG	CAGTCCAGTAATTTTCAAGC	60	
RPE65	6	TTCAAGGGGTAGTGATGACC	GCACAAAATGCTATTCTGAC	60	
RPE65	11	CTTAGGAGCCAAGACTTAAG	GAGGAAACTCAAATGCTACG	60	
PROM1	12	CTCCAGCCTTAGTCCAGCAG	GTCCCATCACAGCAGGATCT	58	
EYS	29	AATCTGCTTCTGGCTTTGTTT	GCCCACTAGCCAGAAAATA	58	
PDE6B	7	CCTGCACACAGACATCCAGT	TGGCAGAGACAAGGAGAAGC	58	
FAM161A	3	TGGTCACATACAACTGAAAGTATAACA	GCTTCTGTTCCTCCCTTGCT	60	
CERKL	8	GGGAACTGAAGAGAAAGTGAGG	GGCTGAGTGAGTAGTTGTTTGC	58	
LRP2	69	TTTTTGCTCCCCATCTTCTG	AATTCAGCAGGTGGGAGTTG	58	
PPIG	14	CATTTCCCCAATTCTTTTTCTT	GATTGTGCGCGTCTATCCTT	58	
CWC22	10	GCTCCTGGAAAGACCAACAG	GCCTCTTTCCAACTAACATGC	58	
ITGA4	24	AAAACCAGGTTGCTTTTTGC	ACCATGCTTGTTCCTTCCAC	58	
PDE1A	7	ACAGGGTGCACTGAAAGTCC	TTCTGGGCAGAGACAGATTG	58	
NCKAP1	11	AGCAACATAAGAAAGGCAGGA	GAAGAATGCAGGGAGGTACG	58	
ZNF804A	4	CCACTGTTGCTGAAGATCCA	TTTGATCTGCTAATGGGACTGA	58	
FSIP2	12	ATCCCTGATTGTGGCAATTC	AAGCAGACTGTTCAAGGGCTA	58	
COL3A1	30	TAGTTCCCACCCAGCTGTTC	CAGCAGCACCCTGAAAATAA	58	
ANKAR	16	TGTCAATATAGCATACCCCTTTG	TTTTTGGCTTATTTCTCCAAGA	58	

Table 1. Sequences of primers and PCR condition.

#### Results

Clinical features. As summarized in Table 2, patient phenotypes included RP and early-onset retinal degeneration (EORD) and STGD (Fig. 1). The clinical presentation of patients in families F2, F4, F5, F7, F8, F10 and F11 showed typical hallmarks of RP symptoms for which affected individuals initially experienced night blindness with progressive visual loss and hemeralopia in the majority of cases. Specifically, affected individual IV-5 (aged 40 years) of F2 family reported poor visual acuity. Funduscopy revealed typical RP changes with macular atrophy in F10 affected members suffering from photophobia; onset of symptoms was reported to occur during the first decade of life, with typical RP changes and normal macula. Fundus examination of patient III-10 (aged 36 years) of F10 revealed normal macular pattern with typical RP, with no nystagmus or hemeralopia. In F11, the index patient V-1 presented typical RP with atrophy of the retina and macular edema.

In addition, clinical data of F9, F12, F13, F14 and F15 (Fig. 2) showed typical hallmarks of RP symptoms for which affected individuals initially experienced night blindness with progressive visual loss and hemeralopia in the majority of cases. Fundoscopy revealed mild optic disc and retinal atrophy without pigmentation. Macula was normal in young patients and showed atrophic alteration in advanced stages.

Intermediate phenotypes between LCA and RP were observed in two families. In F1 and F6 the affected individuals had poor visual acuity at the ages of 10–12 years respectively, with a clinical picture suggesting EORD. Two retinal patterns were identified: pattern 1 presented mid-peripheral deep white dot deposits and virtually no clumped pigmentation (F1), whereas pattern 2 showed mid-peripheral pigmented clumps with no white deposits (F6). Family F3 presented a very advanced form of STGD. Night blindness was the first symptom, followed by intense photophobia. All had useful vision until the second decade of life. Fundus examination revealed atrophy of the macula, pallor of the optic disk and bone spicule-shaped pigment deposits in the periphery. SD-OCT show diffuse atrophy with Epiretinal membrane. Only family F3 presented a Stargardt phenotype. All cases presented non-syndromic phenotype (no associated extraocular abnormalities).

**Mutation analysis.** Following the clinical diagnosis of retinal dystrophy, WES or IROme was performed to screen for all known genes and to look for novel candidate genes. Assuming identity-by-descent in these consanguineous families, we identified and examined autozygous regions when they spanned more than 5 Mb of genomic DNA. After all filtering steps, a few strong candidate genes remained in the autozygous regions of interest. Known RD genes were selected first for direct sequencing.

Mutation analysis in our patients revealed seven novel homozygous mutations. The first mutation found was a missense substitution, c.2293C > T, in exon 23 of CNGB1 (NM\_001297.4), resulting in p.R765C (Fig. 3A) detected in family F2. It is considered to be probably damaging with a score of 1.0 by polyphen-2 and R765 is conserved from mammals to Danio rerio (Zebrafish). The second variant was the substitution of A to G at nucleotide position 1010 in exon 7 of the PDE6B (NM\_001145291.1) and resulted in the substitution of histidine (H) by arginine (R) at codon 337 (p.H337R) (Fig. 3D) in family F10. This variant is predicted to be benign with polyphen-2 (Table 3). This amino acid is also conserved from mammals to Danio rerio. These two mutations were amongst those included in the evaluation of IROme<sup>8</sup>.

Family	F1	F2	F3	F4	F5	F6	F7	F8	F9	F10	F11	F12	F13	F14	F15
Individual	VI-1	IV-5	IV-1	IV-1	IV-5	IV-2	V-1	III-3	IV-2	III-9	V-1	VI-1	V-5	III-5	II-5
Gender	Male	Female	Female	Male	Male	Male	Male	Male	Male	Male	Female	Male	Male	Male	Female
Age (Years)	12	40	35	28	46	10	14	41	38	36	35	14	28	27	38
Age of onset	5	6	12	4	Infancy	Since birth	Since birth	10	22	4	12	4	13	25	Infancy
Age at last follow-up	11	34	31	26	45	9	12	39	37	34	34	14	28	27	38
Legally blind	-	-	+	-	+	+	+	+	+	-	-	-	-	+	-
Photophobia	+	+	+	+	-	+	+	-	+	+++	-	-	+	+	+
Nystagmus	-	-	-	-	-	-	++	++	++	-	-	-	-	-	-
Hemeralopia	-	++	-	++	+	++		+	+	-	++	-	++	+	++
Visual acuity RE/LE	7/10 7/10	2/10 2/10	1/16 1/16	5/10 4/10	1/50 1/50	1/10 1/10	<1/10 <1/10	1/20 1/20	1/10 1/10	3/10 3/10	2/10 2,5/10	4/10 5/10	4/10 2/10	1/10 1/10	2/10 3/10
Fundus photography	mid- peripheral deep white dot deposits and virtually no clumped pigmentatio n	Typical RP changes with bone spicule-shaped pigment deposits in the mid periphery along with atrophy of the retina, narrowing of the waxy optic discs and macutar atrophy	Macular atrophy with pigmentations and peripheral RPE atrophy.	Few bone spicule- shaped pigment deposite the smit deposite along with atrophy of the periphery retina, narrowing of the vessels	Typical RP changes with bone spicule-shaped pigment deposits in the mid periphent along with atrophy of the retina, narrowig of the vessels, atrophy of the macula and slight optic discs	Diffuse retinal atrophy with mid-peripheral pigmented clumps without any white deposits	White deposits in the mid periphery, Mid optic atrophy, Narrowing of the vessels	Macular atrophy with pigmentation s Typical RP-changes with bone spicule-shaped pigment deposits in the mid periphery along with atrophy of the retina, narrowing of the vessels and pallor disc	Diffuse retinal atrophy with pigment clumping and bone spicule pigmentation, attenuated blood vessels with mild optic atrophy	Typical RP changes with few bone spicule-shaped pigment deposits in the mid periphery along with atrophy of the retina, marrowing of the wessels a and slight pallor disc	Typical RP change s with few bone spicule-shaped pigment deposit s in the mid peripher atrophy of the retina, narrowing of these s a and slight pallor disc. Macular	mid- periph eral retinal atroph y and virtuall y no clumpe d pigmen tation mild optic disc atroph y	atrophy of the periphery retina and virtually no clumped pigmentat ion mild optic disc atrophy	atrophy of the periphery retina and virtually no clumped pigmentat ion mild optic disc atrophy	Diffuse retinal atrophy with few pigment dumping pigmentat ion in the peripheral attenuate d blood vessels with mild optic atrophy
coherence tomography		atrophy	atrophy	edema	atrophy	edema	atrophy	atrophy	atrophy		edema			atrophy	atrophy
Other Disease	No	No	No	No	No	No	No	No	No	No	High myopia	No	No	No	No

**Table 2.** Clinical features of the patients examined. + and - symbols indicate presence/absence, as well as degree of a given feature (+ mild, ++ moderate, +++ severe). F: family; RE: right eye; LE: Left eye.

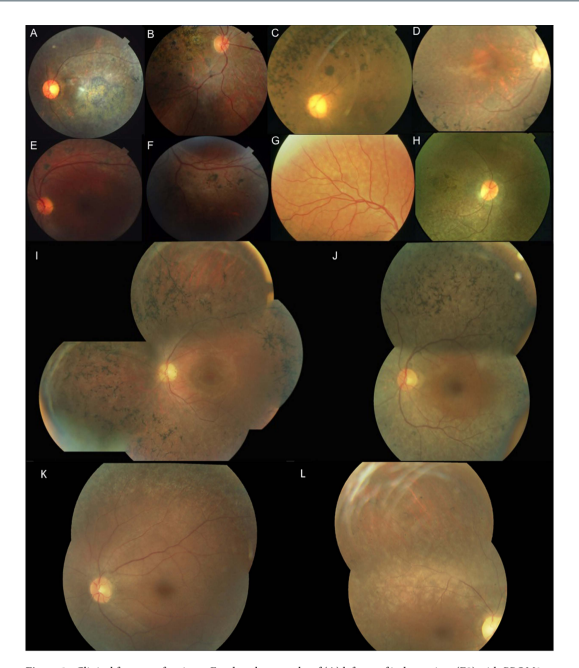
Two new mutations were found in RPE65. The first was in the splice site of RPE65 at position c.1129-2 and changed the canonical splice site second nucleotide to G in family F6 (Fig. 3B). The second mutation was identified in family F7 and was a new compound heterozygous mutations c.[325C > T];[569T > A] resulting in p.(R91W);(V172D) (Fig. 3C). The V172D mutation was not described in ExAC Browser<sup>9</sup>, 1000 genomes<sup>11</sup>, and HGMD<sup>10</sup> databases; this mutation is predicted to be possibly damaging with a score of 0.721. In family F11 we identified several genomic homozygous regions, one of which included FAM161A. Sequence analysis and validation by Sanger sequencing showed a new homozygous deletion of four nucleotides GAAG in exon 3 (c.678\_681delGAAG) (Fig. 3E) resulting in a frameshift and a termination codon at position p.K227Nfs\*17. To our knowledge, this mutation has not previously been reported in the literature or as variation in databases such as 1000 genomes<sup>11</sup> and ExAC. The last mutation was a 4-bp deletion, c.1133 + 3\_1133 + 6delAAGT, located in the donor splice site of intron 8 of CERKL in family F12 (Fig. 3F). This new deletion was present at a homozygous state in the two affected sons and was heterozygous in the parents. It was not present in the unaffected sister. Furthermore, this deletion affects the invariant acceptor site of exon 8, which has not previously been linked to this particular RP phenotype. Minigene analysis confirmed that this variant has a strong impact on the normal splicing pattern of CERKL. Transfection of HEK 293 cells with minigene constructs supporting this change and its wild-type counterpart revealed that c.1133 + 3\_1133 + 6delAAGT did affect the canonical splicing of exon 8 by knocking down its natural 3' splice site (Fig. 4).

As the fundus of affected F12 family members showed atrophy of the peripheral retina with no clumped pigmentation, we looked for this specific phenotype in families with simplex cases or where consanguinity was not evident. Fourteen additional Tunisian families were screened for this specific mutation. This allowed us to identify the homozygous c.1133 + 3\_1133 + 6delAAGT deletion in three seemingly unrelated families (F13, F14, F15), but originating from the same geographical area. Genotype analysis of these three families showed a common homozygous region of 5.7 cM. This is indicative of an unknown consanguinity of these families (Table 4).

The mutations identified in the other families have already been reported in the literature. These changes were two nonsense substitutions: p.R203\* in *CNGB3* (NM\_019098.4) in family F1 and p.W782\* in *ABCA4* (NM\_000350.2) in family F3; two missense mutations: p.R311Q in *NR2E3* (NM\_014249.3) in F4 and p.H182Y in *RPE65* (NM\_000329.2) in F5, one frameshift mutation: c.1354dupT in *PROM1* (NM\_001145848.1) in F8, and one splice site mutation: c.5928-2A > G in *EYS* (NM\_001292009.1) in F9. A summary of these results is presented in Table 3. Cosegregation analysis in each family showed that all affected individuals were homozygotes and unaffected individuals were either heterozygous carriers or homozygous for the wild type allele (Fig. 5).

#### Discussion

Molecular diagnosis of RD is a challenging task given the high genetic heterogeneity of this group of diseases. Aiming to identify the molecular origin of RD in our Tunisian population, we performed molecular analysis of index cases from Tunisian families and revealed a great genetic variety among this population. Ten different genes were involved in 15 families. Previous studies in the Tunisian population have shown a high genetic heterogeneity in RP patients<sup>14</sup> as well as in patients with Usher syndrome<sup>15</sup>. This could be explained by the degree of consanguinity observed in various parts of the country. The genes identified play different roles in the retina. Most genes mutated in this sub-cohort are a part of three major processes in rod photoreceptors and RPE. The first



**Figure 1.** Clinical features of patients. Fundus photographs of (**A**) left eye of index patient (F8) with *PROM1* mutation; (**B**) right eye of IV-4 patient (F3) with *ABCA4* mutation; (**C**) left eye of index patient (F5) with *RPE65* mutation; (**D**) right eye of the index case (F2) with *CNGB1* mutation; (**E**,**F**) left eye of subject IV-1 in family F4 with *NR2E3* mutation; (**G**) right eye of subject VI-1 (F1) with *CNGB3* mutation; (**H**) right eye of IV-1 patient (F6) with *RPE65* mutation; (**I**) left eye of index case IV-2 (F9) with *EYS* mutation, (**J**) left eye of subject III-9 (F10) with *PDE6B* mutation, (**K**) left eye of the proband V-1 (F7) with *RPE65* mutation, (**L**) right eye of subject V-1 (F11) with deletion in *FAM161A*.

three genes, PDE6B, CNGB1 and CNGB3, are implicated in the phototransduction cascade: PDE6B, encoding the beta-subunit of rod cGMP-phosphodiesterase (cGMP-PDE), is a key enzyme of the retinal rod phototransduction cascade. PDE6B was reported by several groups as one of the causative gene associated with arRP, including previous studies of consanguineous Tunisian families<sup>16–19</sup>. Mutations in CNGB1 encoding the beta-subunit of the rod cGMP-gated channel explain about 4% of autosomal recessive non-syndromic RP<sup>4</sup>. We discovered a new p.R765C homozygous missense substitution in exon 23 of CNGB1 in F2. This mutation occurred at an evolutionarily conserved position from mammals to zebrafish and is predicted to damage the protein. So far, only eight point mutations were described with convincing evidence. Four of them were non-sense. Our result concurred with that of Bariel  $et\ al.$ , who found mutation in CNGB1 in a consanguineous French family affected with severe autosomal recessive RP (arRP)<sup>20</sup>. CNGB3 encodes the  $\beta$ -subunit of the cone photoreceptor cGMP-gated channel, essential for phototransduction in all three classes of cones<sup>21</sup>. Mutations in this gene are a common cause of

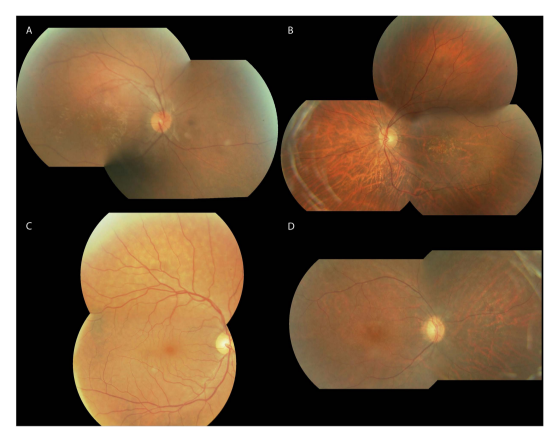


Figure 2. Spectrum of fundus findings among different families (F12, F13, F14, F15) with mutation in CERKL.

achromatopsia<sup>14</sup>. Missense mutations in *CNGB3* have previously been reported in two individuals with conerod dystrophy and in a single individual with a progressive cone dystrophy phenotype<sup>22</sup>. Although the p.R203\* mutation was previously associated with achromatopsia<sup>21</sup>, in family F1 it induced a more severe phenotype that was classified as EORD. In the past, CNGB3 was mainly screened in patients with achromatopsia in which it represents a common cause. With current high-throughput sequencing, most approaches analyze a large set of genes. This non-biased approach will most likely widen the genotype-phenotype correlation of most diseases<sup>23–25</sup>.

We also observed four families with homozygous mutations in the retinoid cycle genes. In ABCA4, a member of the ABC transporter superfamily associated with STGD disease<sup>26</sup>, CRD, RP<sup>27,28</sup> as well as age-related macular degeneration<sup>29</sup>, a p.W782\* was identified in family 3 with STGD phenotype. In RPE65, a vitamin A trans-cis isomerase, a p.H182Y was observed in family 5. This mutation was first reported by Morimura et al. as a compound heterozygous mutations in a single family with two boys affected with LCA30. Interestingly, this amino acid has also been mutated into Asparagine and Arginine as reported in Hanein et al. and Jacobson et al., respectively<sup>31,32</sup>. The p.H182N and p.H182R were associated with LCA<sup>31,32</sup>. Unfortunately, there are not enough families with mutations at H182 to establish a strong genotype-phenotype correlation for this position. Mutation analysis in the region of homozygosity in family F6 revealed a new mutation c.1129-2A > G that affects the consensus splice site of exon 11 of RPE65 and segregates with EORD. A compound heterozygous mutation p.(R91W);(V172D) was discovered in one family (F7). To date, a number of different RPE65 mutations have been reported in patients with retinal dystrophies classified as LCA, autosomal recessive EORD or arRP<sup>30,33,34</sup>. However, in our cohort we found three different RPE65 mutations in three different families that correlated with typical RP phenotype despite the early age of onset. Morimura et al., reported a family in which a child with LCA phenotype is the offspring of two parents with RP<sup>23</sup>. A recent Chinese study, including 88 candidate genes in 179 families with RD, confirmed that the average age at onset was 5 years in patients carrying mutation in RPE65 gene<sup>24</sup>. In patients from family F4 harboring the p.R311Q, the most prevalent mutation in the nuclear receptor subfamily 2, group E, member 3 (NR2E3), we observed typical RP with macular atrophy, as already reported35. Nevertheless, Gerber et al. have shown that the p.R311Q mutation could cause late-onset arRP in Crypto-Jewish population<sup>36</sup>. NR2E3 recessive mutations are usually associated with enhanced S-cone syndrome, an autosomal recessive retinopathy in which patients have increased sensitivity to the perception of blue light while the dominant p.G56R mutation is associated with severe  $RP^{21,22}$ . A homozygous splice alteration in EYS (c.5928-2A > G) was reported in family F823. EYS is likely to play a role in the modeling of retinal architecture37. In previous studies, various mutations including point mutations, splice-site mutations and gross rearrangements were identified in patients with ar  $RP^{23,38-40}$  and cone rod dystrophies  $^{41}$ . The phenotype observed in family F8 resembles the RP phenotype reported by Barragan et al.<sup>38</sup> in the Spanish population. The advanced age of onset of the disease could be correlated with our molecular result, as Huang et al. found a relationship between the advanced average age

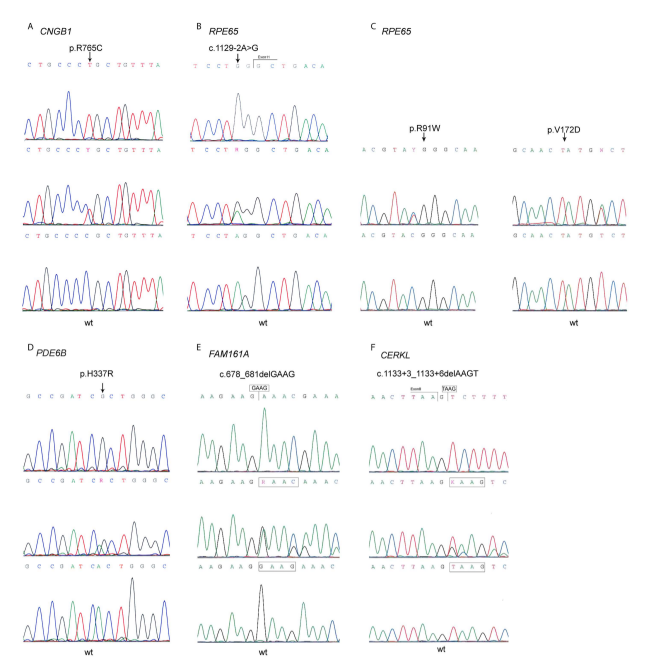


Figure 3. Chromatograms of the six novel mutations in *CNGB1*, *RPE65*, *PDE6B*, *FAM161A* and *CERKL*. For each mutation, wild-type (wt) sequence is depicted with the homozygous and heterozygous sequences. (A) c.2293C > T; p.R765C (*CNGB1*), (B) c.1129-2A > G (*RPE65*), (C) c.[325C > T];[569T > A] (*RPE65*), (D) c.1010A > G; p.H337R (*PDE6B*), (E) c.678\_681delGAAG (*FAM161A*), and (F) c.1133 + 3\_1133 + 6delAAGT(*CERKL*). Arrows indicate the position of the mutated nucleotide.

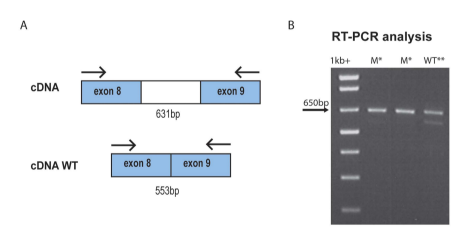
at onset and EYS mutation<sup>24</sup>. The mutation in EYS appears to be a frequent cause of arRP (15.9%) in the Spanish population<sup>28</sup>. Unfortunately, the prevalence of EYS mutations remains to be established in Tunisian population. Our preliminary analysis of the mutation load in the Tunisian population does not allow us to draw a similar conclusion. The fifteen families described here represent a selection that underwent whole-exome sequencing and thus do not represent a correct estimation of the yield of this approach.

In addition, a previously described variant (c.1354dupT) in *PROM1*, which encodes a pentaspan transmembrane glycoprotein that maintains the structural integrity of retina, was observed in patient III:1 from family F7 affected with RP<sup>42</sup>. Mutations in *PROM1* are usually associated with RP or CRD, which is the case in this family<sup>43</sup>.

We also report the identification of a new small deletion in exon 3 of *FAM161A*, a ciliary gene previously associated with RP<sup>44</sup>. This mutation causes a shift in the reading frame resulting in a premature termination codon at position p.K227Nfs\*17, presumably inhibiting protein production due to the action of the nonsense mediated mRNA decay. Most currently known mutations<sup>44,45</sup> cluster in exon3 of *FAM161A*, which is by far the largest coding exon of this gene. FAM161A is member of the growing list of ciliary proteins implicated in human diseases.

Family ID	Disease	Genotyping Method	Size of homozygous region, in Mb	Chromosome	Gene	DNA mutation	Predicted protein variant	Coverage	Reference sequence	Previously reported	SIFT	Polyphen
F1	Eord	WES	30.4	8q21.3	CNGB3	c.[607C > T];[607C > T]	p.(R203*); (R203*)	26	NM_019098.4	[21]	_	_
F2	RP	IROme	_	16q21	CNGB1	c.[2293C > T];[2293C > T]	p.(R765C); (R765C)	_	NM_001297.4	This study and [8]	0	1,00
F3	Stargardt	WES	5.8	1p22.1	ABCA4	c.[2345G > A];[2345G > A]	p.(W782*); (W782*)	5	NM_000350.2	[28]	_	_
F4	RP	WES	12.2	15q23	NR2E3	c.[932G > A];[932G > A]	p.(R311Q); (R311Q)	24	NM_014249.3	[35]	0.10	0.627
F5	RP	WES	54.1	1p31.3-p31.2	RPE65	c.[544C>T];[544C>T]	p.(H182Y); (H182Y)	36	NM_000329.2	[30]	0	0.872
F6	Eord	WES	18.2	1p31.3-p31.2	RPE65	c.[1129-2A > G]; [1129-2A > G]	_	30	NM_000329.2	This study	_	_
F7	RP	IROme	_	1p31.3-p31.2	RPE65	c.[325C > T];[569T > A]	p.(R91W); (V172D)	_	NM_000329.2	This study	0.01/0.01	0.997/0.721
F8	RP	WES	9	4p15.32	PROM1	c.[1354dupT];[1354dupT]	p.(Y452fs*12); (Y452fs*12)	12	NM_001145848.1	[42]	_	_
F9	LCA	WES	58	6q12	EYS	c.[5928-2A > G]; [5928-2A > G]	_	24	NM_001292009.1	[23]	_	_
F10	RP	IROme	_	4p16.3	PDE6B	c.[1010A > G];[1010A > G]	p.(H337R); (H337R)	_	NM_001145291.1	This study and [8]	1.00	0.046
F11	RP	WES	13	2p15	FAM161A	c.[678_681delGAAG]; [678_681delGAAG]	p.(K227Nfs*17); (K227Nfs*17)	94	NM_001201543	This study	_	_
F12	RP	WES	8	2q31.3	CERKL	c.[1133+3-1133+6delAAGT]; [1133+3-1133+6delAAGT]	_	21	NM_201548.4	This study	_	_
F13	RP	_	_	2q31.3	CERKL	c.[1133+3-1133+6delAAGT]; [1133+3-1133+6delAAGT]	_	_	NM_201548.4	This study	_	_
F14	RP	_	_	2q31.3	CERKL	c.[1133+3-1133+6delAAGT]; [1133+3-1133+6delAAGT]	_	_	NM_201548.4	This study	_	_
F15	RP	_	_	2q31.3	CERKL	c.[1133+3-1133+6delAAGT]; [1133+3-1133+6delAAGT]	_	_	NM_201548.4	This study	_	_

Table 3. Homozygous regions and mutations identified in this study.



**Figure 4.** RT-PCR products with primers in exon 8 and 9. (A) Schematic representation of normally spliced mRNA, (B) Aberrantly spliced mRNA. In the gel electrophoresis we can observe absence of splicing in the mutated patient (column M). In the WT sequence (column WT), splicing was not complete. \*Non-spliced fragment, \*\*spliced fragment.

Phenotypes of ciliopathies are quite diverse and may involve impairment of multiple organs or functions such as kidney, brain, bones, obesity and vision<sup>46</sup>. However, in the patients investigated in this study we did not observe any other obvious features or anamnestic history of extraocular symptoms typically related to ciliopathies. The phenotype was typical of RP.

Finally, it is of note that the analysis of four families (F13, F14, F15 and F16) originating from the same Tunisian region revealed a new homozygous deletion c.1133 + 3\_1133 + 6delAAGT in the Ceramide kinase Like (CERKL) gene in association with RP without pigmentation. Mutations in CERKL have been described in RP (RP26) together with significant macular involvement during the early stages of the disease<sup>47</sup> and cone-rod dystrophy which progresses to an RP-like phenotype in advanced stage<sup>48</sup>. However, the exact function of CERKL

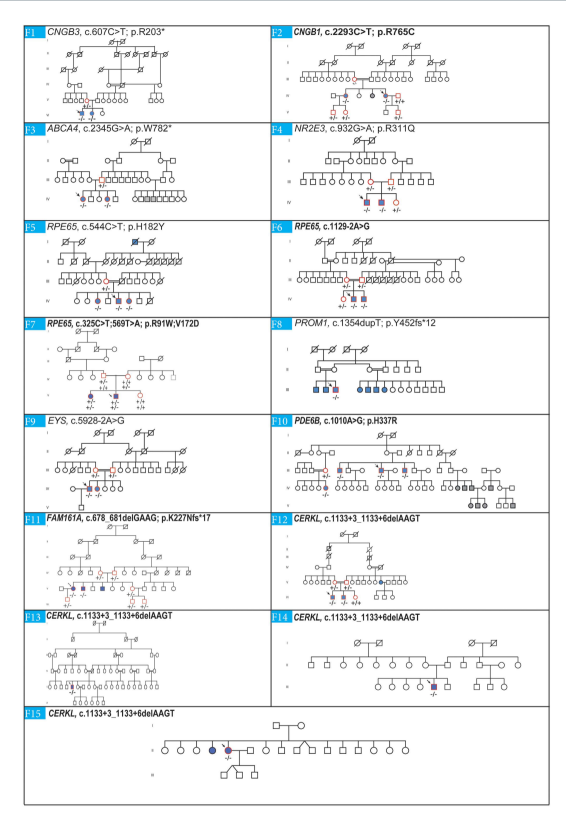


Figure 5. Overview of the pedigree structure of the families and segregation analysis of disease causing variants in the RD cohort. Arrows point to the probands. Affected individuals are indicated with filled symbols (blue), suspected (grey), whereas unaffected relatives are indicated by open symbols. +: wild type allele; -: mutation. All mutations are homozygous except for family F7.Genes with new mutations are in bold.

remains unknown. Genotype analysis of these four families showed a common disease-associated haplotype and supports the hypothesis of a common ancestor in this area.

rs.	Distance position	F12 (VI-1)	F13 (V-5)	F14 (III-5)	F15 (II-5)
rs4667591	-12.4 M	A/C	C/C	A/C	A/C
rs1050354	-11.9 M	T/A	A/A	T/A	T/A
rs4894140	-1.5 M	C/C	C/C	C/C	C/C
rs1143676	-0.18 M	A/A	A/A	A/A	A/A
CERKL	0	_	_	_	_
rs3754929	+0.67 M	T/T	T/T	T/T	T/T
rs1400130	+1.43 M	G/G	G/G	G/G	G/G
rs12476147	+3.3 M	T/T	T/T	T/T	T/T
rs17228441	+4.2 M	C/C	C/C	C/C	C/C
rs1800255	+7.4 M	G/A	G/A	G/A	G/A
rs1225090	+8.1 M	G/T	T/T	G/T	G/T

Table 4. Common homozygous genomic region.

Despite intensive research and studies, it is still very difficult to screen for specific genes based on clinical observation and severity of disease. Molecular diagnosis of RD is a challenging task given the important genetic heterogeneity of this group of diseases.

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#### **Author Contributions**

D.F.S. and H.I. identified the mutations; A.C., Y.F., F.K. and L.E.M. referred patients and clinical data; D.F.S. and H.I. wrote the paper; D.F.S., A.C., Y.F. and H.I. prepared the figures; H.I. and N.A.P conducted the minigene experiments; D.F.S. and L.E.M designed the experiments. All authors reviewed and approved the manuscript.

#### **Additional Information**

**Competing financial interests:** The authors declare no competing financial interests.

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## **OPEN Corrigendum: Identifying** mutations in Tunisian families with retinal dystrophy

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This Article contains errors. The position of the mutation p.(R91W); (V172D) was incorrectly calculated, taking as a starting point the beginning of cDNA rather than the start codon. The correct position of the mutation is c.[271C>T]; [515T>A]. As a result of this the following changes in the Article are made:

The legend of Figure 3 is incorrect,

"(C) c.[325C > T]; [569T > A] (RPE65)"

should read:

"(C) c.[271C > T]; [515T > A] (RPE65)".

In Table 3 for the family ID F7, the DNA mutation should read 'c.[271C > T]; [515T > A]'.

In Figure 5 for family F7 'RPE65, c.325C > T; 569T > A; p.R91W; V172D' should read 'RPE65, c.271C > T; 515T > A; p.R91W; V172D'. The correct Figure 5 appears below as Figure 1.

In the Results section under the subheading 'Mutation analysis',

"The second mutation was identified in family F7 and was a new compound heterozygous mutations c.[325C>T]; [569T > A] resulting in p.(R91W); (V172D) (Fig. 3C)".

should read:

"The second mutation was identified in family F7 and was a new compound heterozygous mutations c. [271C > T]; [515T > A] resulting in p.(R91W); (V172D) (Fig. 3C)".

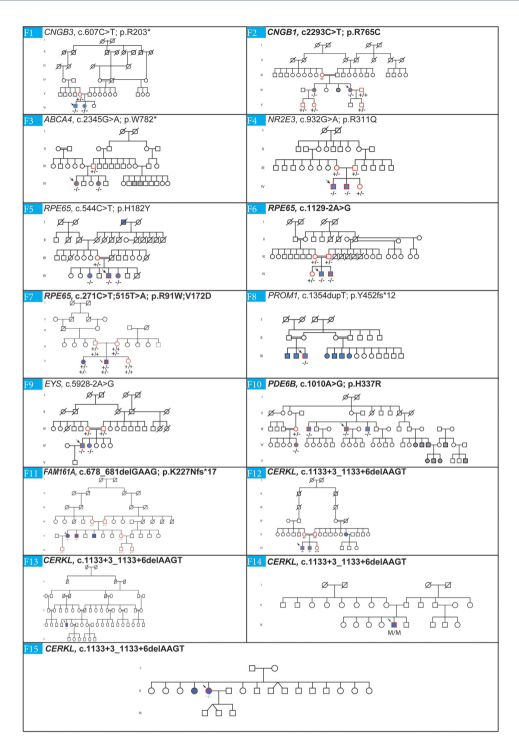


Figure 1.

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