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#### ORIGINAL ARTICLE

## Novel insights into the molecular pathogenesis of CYP4V2associated Bietti's retinal dystrophy

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#### Keywords

Bietti, crystalline dystrophy, CYP4V2, retinal dystrophy

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#### Abstract

Bietti's crystalline dystrophy (BCD) is a rare, autosomal recessive retinal degenerative disease associated with mutations in CYP4V2. In this study, we describe the genetic and clinical findings in 19 unrelated BCD patients recruited from five international retinal dystrophy clinics. Patients underwent ophthalmic examinations and were screened for CYP4V2 mutations by Sanger sequencing and quantitative polymerase chain reaction (qPCR) copy number variation screening. Eight CYP4V2 mutations were found in 10/19 patients, including three patients in whom only monoallelic mutations were detected. Four novel mutations were identified: c.604G>A; p.(Glu202Lys), c.242C>G; p.(Thr81Arg), c.604+4A>G; p.(?), and c.1249dup; p.(Thr417Asnfs\*2). In addition, we identified a heterozygous paternally inherited genomic deletion of at least 3.8 Mb, encompassing the complete CYP4V2 gene and several other genes, which is novel. Clinically, patients demonstrated phenotypic variability, predominantly showing choroidal sclerosis, attenuated vessels, and crystalline deposits of varying degrees of severity. To our knowledge, our study reports the first heterozygous CYP4V2 deletion and hence a novel mutational mechanism underlying BCD. Our results emphasize the importance of copy number screening in BCD. Finally, the identification of CYP4V2-negative patients with indistinguishable phenotypes from CYP4V2-positive patients might suggest the presence of mutations outside the coding regions of CYP4V2, or locus heterogeneity, which is unreported so far.

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#### Introduction

Bietti crystalline dystrophy (BCD) (OMIM 210370) is a rare retinal degenerative disease that is inherited in an autosomal recessive pattern (Li et al. 2004). First described in three patients by the Italian ophthalmologist Bietti (1937) in Rome, BCD is defined and characterized by glistening crystalline deposits in the fundus associated with atrophy of the retinal pigment epithelium (RPE) and choroidal sclerosis (Bietti 1937; Rossi et al. 2013). Crystal deposits at the corneal limbus have also been documented (Rossi et al. 2013). Clinically, patients affected with BCD present between the second and fourth decade of life with impaired vision, nyctalopia, and paracentral scotomas. Vision loss and concentric visual field constriction continue progressively, often resulting in severe visual impairment by the fifth or sixth decade (Li et al. 2004; Lee et al. 2005).

BCD has been reported to be more prevalent in Asian populations, although patients of European, Middle Eastern, African, and North and South American origin have also been documented (Hu 1987; Li et al. 2004). Moreover, it has been estimated that up to 3% of patients initially diagnosed with nonsyndromic retinitis pigmentosa can be accounted for by BCD (Mataftsi et al. 2004). BCD is known to be caused by mutations in the CYP4V2 gene, localized on chromosome 4q35 (Li et al. 2004). Currently, up to 57 mutations in CYP4V2 have been associated with BCD; the majority are missense/nonsense mutations, as well as several small insertions, deletions, and splicing mutations (Li et al. 2004; Lee et al. 2005; Lin et al. 2005; Shan et al. 2005; Wada et al. 2005; Jin et al. 2006; Lai et al. 2007; Zenteno et al. 2008; Mamatha et al. 2011; Xiao et al. 2011; Yokoi et al. 2011; Haddad et al. 2012; Manzouri et al. 2012; Parravano et al. 2012; Song et al. 2013; Halford et al. 2014; Yin et al. 2014). CYP4V2 encodes a member of the cytochrome P450 superfamily, characterized as a fatty acid oxidase involved in lipid metabolism (Nakano et al. 2009). Surprisingly, evidence of altered lipid metabolism has also been demonstrated in fibroblasts and lymphocytes of patients with BCD, in which synthesis of n-3 polyunsaturated fatty acids was decreased (Lee et al. 2001). Crystalline deposits have also been reported in these cells (Wilson et al. 1989). Furthermore, patients with BCD have been shown to have abnormal serum fatty acid profiles, with increased stearic acid and reduced oleic acid concentrations (Lai et al. 2010). These findings suggest that BCD represents a systemic condition that is not limited to the eye. However, it is still unclear whether these systemic abnormalities are actually disease-causing or whether they are subclinical (Li et al. 2004). It has been hypothesized that the retinal crystals represent lipids or fatty acids (Lai et al. 2010). Finally, the phenotypic spectrum of *CYP4V2* mutations has been expanded to autosomal recessive retinitis pigmentosa (arRP), without intraretinal crystals being observed in affected patients (Mataftsi et al. 2004).

In this study, we present novel genetic and clinical findings in a large international cohort of 19 probands from unrelated families with apparent Bietti crystalline retinal dystrophy.

#### **Materials and Methods**

#### Subject enrolment and clinical evaluation

Nineteen patients were recruited from and evaluated at the McGill Ocular Genetics Laboratory and Clinic at the McGill University Health Centre, in Montreal Canada; the Centre for Ophthalmology at the University of Tübingen, in Germany; the Ophthalmic Genetics Clinic at the Ghent University Hospital, in Belgium; the Rare Retinal Disease Centre (Siegburg Eve Center), Germany; and INSERM in Montpellier, France. Informed consent was obtained and research protocols adhered to the tenets of the Declaration of Helsinki. Institutional Review Board (IRB)/Ethics Committee approval was obtained. Patients had histories and pedigree analysis taken, and underwent ophthalmic examination including visual acuity (VA) testing, Goldmann visual field testing, refraction, slit lamp examination, dilated fundus examination, fundus photography, fundus autofluorescence (FAF), optical coherence tomography (OCT), and full-field flash electroretinography (ERG), recorded in accordance with the guidelines of the International Society for Clinical Electrophysiology of Vision (Marmor et al. 2009). One patient did not have Goldmann visual field testing nor OCT testing. In addition, fluorescein angiography was also performed in two patients. Phlebotomy in ethylenediaminetetraacetic acid tubes was performed to collect venous blood for genetic studies.

#### **Mutation analysis**

# Homozygosity mapping (only performed in Lebanese family A)

Genomic DNA was isolated from lymphocytes by standard salting out procedures (Miller et al. 1988). DNA samples of all four affected individuals from family A were genotyped on the GeneChip Genome-Wide Human SNP Array 5.0 that contains 500,000 polymorphic SNPs in addition to 420,000 nonpolymorphic probes for the detection of germline copy number variations (CNVs) (Affymetrix, Santa Clara, CA). Array experiments were performed according to protocols provided by the manufacturer. The 5.0 array data were genotyped using Affymetrix Genotype Console (version 2.1), subsequently regions of homozygosity were identified using Partek Genomics Solution (version 6.1), as described previously (Collin et al. 2011). Regions containing more than 250 consecutive homozygous SNPs were considered as homozygous regions, on average corresponding to a genomic size of 1 Mb or more.

#### **Mutation analysis**

All exons and intron-exon boundaries of *CYP4V2* (NM\_207352) were amplified under standard polymerase chain reaction (PCR) conditions using primers listed in

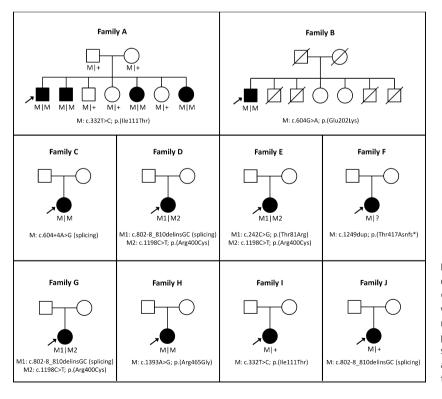
Table S1. PCR products were purified on Nucleospin Plasmid Quick Pure columns (Machery Nagel, Düren, Germany) and sequenced in sense and antisense directions with dye termination chemistry on a 3730 or 2100 DNA analyzer (Applied Biosystems, Carlsbad, CA).

# Bioinformatic analysis and evolutionary comparison for missense mutations

For each of the missense changes identified in this study, the potential pathogenicity was assessed using online prediction software tools SIFT (Sorting Intolerant from Tolerant) and PolyPhen (Ng and Henikoff 2003). Grantham and PhyloP scores were also determined. In addition, the Exome Variant Server database was checked for the presence and minor allele frequencies of these novel mutations. Four computational programs, SpliceSite finder-like, MaxEntScan (Yeo and Burge 2004), NNSPLICE (Reese et al. 1997), and Human Splicing Finder (Desmet et al. 2009) were employed to predict the effect on the canonical acceptor and donor splice sites.

#### CNV screening using qPCR and SNP chip arrays

CNV screening on genomic DNA was performed in the families with only one *CYP4V2* mutation (probands F, I, and J), using 13 quantitative PCR (qPCR) assays, covering



**Figure 1.** Pedigrees of 10 families with Bietti crystalline dystropy carrying mutations in *CYP4V2*. Affected individuals are indicated with filled symbols, whereas unaffected relatives are indicated by open symbols. The probands are indicated with an arrow and slashed symbols indicate deceased. Mutated alleles are indicated with M symbols and wild-type allele with plus symbols.

Table 1. Mutation table.

Proband	Туре	Allele 1		Allele 2	
A	Homozygous	c.332T>C	p.lle111Thr	c.332T>C	p.lle111Thr
В	Homozygous	c.604G>A	p.Glu202Lys	c.604G>A	p.Glu202Lys
С	Homozygous	c.604+4A>G	splicing	c.604+4A>G	Splicing
D	Compound heterozygous	c.802-8_810delinsGC	Splicing	c.1198C>T	p.Arg400Cys
E	Compound heterozygous	c.242C>G	p.Thr81Arg	c.1198C>T	p.Arg400Cys
F	Compound heterozygous	c.1249dup	p.Thr417Asnfs*2	Genomic deletion	Genomic deletion
G	Compound heterozygous	c.802-8_810delinsGC	Splicing	c.1198C>T	p.Arg400Cys
Н	Homozygous	c.1393A>G	p.Arg465Gly	c.1393A>G	p.Arg465Gly
I	Heterozygous	c.332T>C	p.lle111Thr	No second	allele found
J	Heterozygous	c.802-8_810delinsGC	splicing	No second	allele found

the 11 exons of *CYP4V2* and two reference genes (*ZNF80* and *GPR15*) on the LightCycler 480 (Roche, Basel, Switzerland). Four controls were included in each experiment. Conditions and primers can be found in Table S2. Data analysis was performed using qBasePlus (Biogazelle, Zwijnaarde, Belgium).

Genome-wide SNP chip analysis was performed using the HumanCytoSNP-12 BeadChip platform (Illumina, San Diego, CA) (proband F) in order to delineate the deletion.

#### Results

#### Patient demographics and genotyping

In a consanguineous Lebanese family that was initially diagnosed with atypical RP (Fig. 1A), genome-wide SNP

array analysis was combined with homozygosity mapping to identify genomic regions that could potentially harbor the causative genetic defect. In total, only two homozygous regions were identified that were identical between the four affected siblings, one of which harbored the *CYP4V2* gene, located on chromosome 4. Sequence analysis revealed a homozygous missense mutation in *CYP4V2*, c.332T>C; p. (Ile111Thr) that completely segregated in the family (Fig. 1A). Most notably, the proband presented with severe choroidal sclerosis but no crystals were found on fundus examination. Upon further family examination, an affected younger sibling, however, was noted to have diffuse crystalline deposits in the fundus.

Subsequently, 18 additional patients were ascertained that were diagnosed with retinal dystrophy and presented or had presented with crystalline deposits. Sequence

Table 2. Pathogenicity predictions for missense and splice site mutations based on in silico analyses.

Amino acid ch	nange predictions									
DNA change	Mutation conseque	ience E	Exome variant server <sup>1</sup>	Phylo	Grantham score	SIFT	PolyPhen		Refere	nces
c.242C>G	p.(Thr81Arg)	-	_	2.55	71	Tolerated	Benign		Novel	
c.332T>C	p.(lle111Thr)	-	-	4.56	89	Deleterious	s Probably d	amaging	Li et al	. (2004)
c.604G>A	p.(Glu202Lys)	-	-	5.61	56	Deleterious	s Probably d	amaging	Novel	
c.1198C>T	p.(Arg400Cys)	2	2/13004 MAF 0.0154	4.48	180	Deleterious	s Probably d	amaging	Lai et a	al. (2007)
c.1393A>G	p.(Arg465Gly)	2	2/13004 MAF 0.0077	1.25	125	Deleterious	Probably d	amaging	Rossi e	t al. (2013)
Splicing predic	tion									
	Mutation									
DNA change	consequence F	Position	SSF (0-100)		MaxEnt (0–12)	NNSPL	ICE (0–1)	HSF (0-	100)	Reference
c.604+4A>G	Altered splicing	c.604	84.07 ⇒ 73.99 (-12	2.0%)	8.95 ⇒ 4.29 (-52.1	%) 0.98 =	⇒ 0 (−100%)	92.33 = (-9.0%		Novel

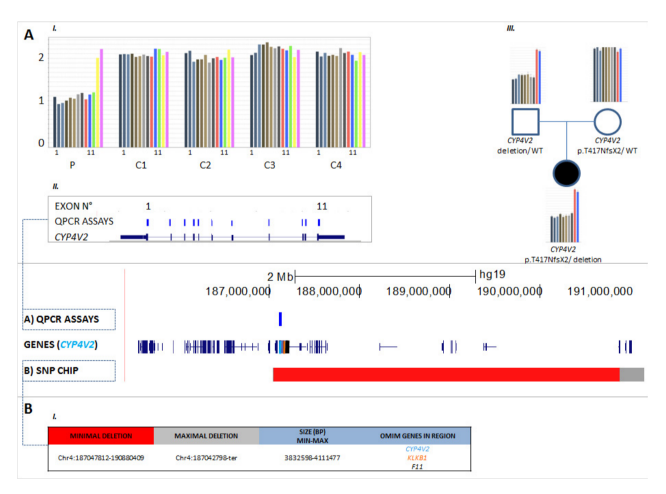
List of missense and splice site mutations identified in this study and predictions of their consequences with the use of in silico program (SIFT and PolyPhen). Splicing prediction shows the percent decrease in comparison to the original splice donor site scores. In addition, HSF predicted a novel splice donor site in position c.604+4. SIFT, sorting intolerant from tolerant; MAF, minor allele frequency; HSF, Human Splicing Finder; SSF, Splice Sequence Finder; NNSPLICE, Splice Site Prediction by Neural Network.

<sup>1</sup>Heterozygous alleles of total number of chromosomes.

analysis of all exons and intron–exon boundaries of *CYP4V2* in this cohort revealed nine more patients with *CYP4V2* mutations, four carrying apparent homozygous mutations, three with compound heterozygous mutations, and two individuals with only one heterozygous intragenic variant (Fig. 1). All these patients were of Caucasian origin.

Patients with *CYP4V2* mutations included seven females and three males, with ages ranging between 33 and 77 years. Initially, eight different mutations in *CYP4V2* were identified, including four previously documented and four novel mutations. The novel mutations were two missense, one splice site and one frameshift mutation; namely c.604G>A; p.(Glu202Lys), c.242C>G; p.(Thr81Arg), c.604+4A>G; p.(?), and c.1249dup; p.(Thr417Asnfs\*2) (Table 1). The novel missense mutations were not observed in the comprehensive Exome Variant Server database, while the already published variants p.(Arg400Cys) and p.(Arg465Gly) have been identified with low minor allele frequencies of 0.0154 and 0.0077, respectively. The most prevalent mutations identified in our cohort were the c.1198C>T; p.(Arg400Cys), c.332T>C; p.(Ile111Thr), and c.802-8\_810delinsGC; p.(?) mutations, each accounting for 3 of the 18 mutant alleles identified in the probands. We found the p. Arg400Cys mutation in three patients, the p. Ile111Thr in two patients, and the c.802-8\_810delinsGC; p.(?) change in three patients.

To evaluate the pathogenicity of the novel mutations, in silico analysis using a variety of prediction programs was performed. The two novel missense mutations, p.(Thr81Arg) and p.(Glu202Lys), affect amino acid residues that are highly conserved among vertebrate species. Both substitutions have a high Grantham score (Table 2). Moreover, the p.(Glu202Lys) mutation, is predicted to be



**Figure 2.** Detection and delineation of a *CYP4V2* deletion in proband F using qPCR and SNP-chip analysis. C (control), P (patient), WT (wild type). (A) I and II: qPCR on proband F revealed a copy number of one for all 11 assays of *CYP4V2*, corresponding with a heterozygous total gene deletion. The same was found for the other deleted genes; KLKB1 and F11. III: Demonstration of paternal origin of the deletion. (B) I: SNP-chip analysis demonstrates deletion with size ~4 Mb, spanning the entire *CYP4V2* gene and several other genes, two of which are OMIM genes (*KLKB1* and *F11*).

					Visual acuity	ty					Visual field			
			Age at		Initial visual acuity	Initial visual acuity	Visual Acuity	Visual Acuity	Refraction	Refraction	Initial visual	Initial visual		Visual field
Family ID	□	Age	onset	Gender	(OD)	(OS)	(OD)	(OS)	(OD)	(OS)	field (OD)	field (OS)	Visual field (OD)	(OS)
A	MOGL	57	30	Σ	N/A	N/A	20/100	20/50	-0.50 +2.00	Plano + 1.50	N/A	N/A	70° (V4e) 65° (III4e)	70° (V4e) 65°
	3254								× 180°	× 180°				(III4e)
а	MOGL	67	45	Σ	N/A	N/A	20/150	12/400	-1.50 +1	-1.50 +1	N/A	N/A	70° with pericentral	70° with
	3138								× 180°	$\times 180^{\circ}$			detect (V4e),	patchy dofects
													peripnerai isiano (14e)	derects (\/4e) 5°
														with
														peripheral island (I4e)
U	13573-	47	30	ш	20/100	20/200	1/30	Ч	-4.75 -1.75	-5.0 - 1.5	Concentric narro	Concentric narrowing :to 5° with	residual islands peripheral (target	eral (target
	BD				(age 38)	(age 38)			× 95°	~ 90°	target III4e, peripheral residual slim islands (age 38)	pheral residual 2 38)	V4e), central island <3°	ŝ
Δ	3338-	54	24	ш	N/A	N/A	МΗ	МH	-1.75 -0.5	-0.75 -1.0	N/A	N/A	Residual islands peripheral (target	eral (target
	BD								$\times$ 76°	× 66°			V4e)	
ш	3549-	41	20	ш	20/20	20/20	20/20	20/20	-1.25 -0.5	-1.0 -0.75	N/A	N/A	Normal outer boundaries with II14e,	ies with III4e,
	BD								× 105°	× 77°			decreased sensitivity and "patchy"	and "patchy"
			č						ī	ī		-	defects with 14e and 13e targets	Ise targets
ш	ΚW	49	, ,	LL_	20/40	20/40	ΣH	20/400	Plano	Plano	Normal	Normal	Mild concentric	Mild
											peripheral	peripheral	constriction,	concentric
											limits,	limits,	central scotoma	constriction,
											pericentral	pericentral	(V4), residual	central
											sensitivity	sensitivity	temporal crescent	scotoma
											loss (I2 not	loss (I2 not	(III4), I2 and I3	with small
											seen),	seen),	not seen	preserved
											central relative	enlarged		patch nasal
											scotoma (I3	blind spot		to blind spot
											and 14),			(V4)
											enlarged hlind snot			
U	10906-	77	57	ш	20/50	20/200	20/100	20/400	+1.0 -2.0	0#	N/A	N/A	N/A	N/A
	BD				(age 75)				× 110°					
т	RCD	38	30	ш	20/30		20/40	20/30	Plano	Plano	N/A	N/A	Mild concentric	Considerable
													constriction,	concentric
													enlarged blind	constriction,
													spot (V4)	enlarged
													considerable sensitivity loss	blind spot
													Incol familiar	/
														(Continued)

Table 3. Clinical features of 10 BCD patients with mutations in CYP4V2.

				Visual acuity	ty					Visual field			
Family	0	Age at Age at	at t Gender	Initial visual acuity (OD)	Initial visual acuity (OS)	Visual Acuity (OD)	Visual Acuity (OS)	Refraction (OD)	Refraction (OS)	Initial visual field (OD)	Initial visual field (OS)	Visual field (OD)	Visual field (OS)
												doughnut shaped annular scotoma (I4), I2 and I3 not seen	considerable sensitivity loss, doughnut shaped annular scotoma (14), 12 and 13 not seen
_	11431- 3 USHII	33 21	ш	20/25 (age 22)	20/40 (age 22)	20/40	20/50	-3.25 -1.25 × 7°	-4.0 -1.5 × 174°	Concentric narrowing to 40° with target III4e (age 22)	ving to 40° (age 22)	Concentric narrowing to 10°, no peripheral islands (target III4e)	g to 10°, no arget III4e)
<b>_</b>		51 18	Σ		20/20	20/40	20/40	Plano	Plano	Concentric reduction, ring scotoma and residual central visual field island of central 5°	Constriction of peripheral visual field, ring scotoma, finally residual central island of central 5°	- MA	AN
	Symptoms				Morp	Morphology							
Family	Nyctalopia	Photophobia		Color vision defect	ect Cornea	lea	Le	Lens	Mã	Macula	Peripheral Retina	Retina	
A	$\succ$	z	Z		Clear	-	Ea	Early cortical changes OS		Relative foveal sparing	Areas of R	Areas of RPE atrophy and choroidal sclerosis, normal vessel caliber peripheral hony spicules	bidal sclerosis, al hony spicules
ß	≻	z	z		Clear	2	S	PSCC	Mâ	Macular involvement	Crystals in choroidal	Crystals in periphery and posterior pole, choroidal sclerosis, tigroid	ior pole,
U	≻	≻	z		Clear	2	Ŭ	Cortical Opacities		Atrophic	Optic disk: RPE atrop	Optic disks vital, narrow vessels, generalized RPE atrophy, crystalline deposits	, generalized ts
Δ	~	z	Z		Limb dep	Limbus: crystalline deposits		Crystalline deposits, PSCC	-	Huge macular hole with neurosensoric detachment on OD, macula atrophic OS	ent : OS	Optic disks vital, narrow vessels, generalized RPE atrophy, crystalline deposits	, generalized ts
ш	~	Z	>		Clear	2	U	Clear	No	None commented		Optic disks vital, peripapillary atrophy, vessels moderately attenuated, posterior pole with patchy RPE-atrophy and crystalline deposits, some pigment clumps	rophy, vessels ior pole with lline deposits,
													(Continued)

Table 3. Continued.

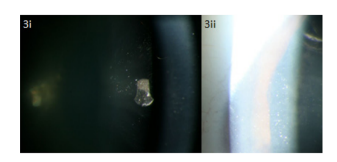
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Endity         National         Constant of the constantont of the constant of the c		Symptoms			Morphology				
Y         N         Y (severe RG, severe BY)         Very stall performant         Mild lemicular scleosis         Extreme outer retinal and crystals         Patry outer crystals           Y         Y (mid)         NA         Clear         Peudophakic         Severe atopy, crystals and grigment para very findity         Performant         Perdophakic         Severe atopy, crystals and grigment para very findity         Performant         Performant<	Family	Nyctalopia	Photophobia	Color vision defect	Cornea	Lens	Macula	Periph	eral Retina
Y     Y (mild)     NA     Clear     Severe atrophy, crystals and figmentary value mice. Severe atrophy, crystals and figmentary value mice fields.       Y     N     Y (medium R/G     Clear     Evelophatic     Severe atrophy, crystals and figmentary value mice fields.       Y     Y     Y     Y     Outor ential and fields.     Partity outer ential and fields.       Y     Y     Y     Y     Clear     Control opacities     Macular etema and inferior.     Opic dists.       Y     Y     Y     Y     Control opacities     Macular etema and inferior.     Opic dists.       Y     Y     Y     Control opacities     Macular etema and inferior.     Opic dists.       Y     Y     Y     Control opacities     Macular etema and inferior.     Opic dists.       Y     Y     Y     Constance     Macular etema and inferior.     Opic dists.       Cons: 55% residual function     Cons: 55% residual function     Opic dists.     Opic dists.       Cons: 50% residual function     Cons: 50% residual function     Opic dists.     Opic dists.       Non recordable     Found definance in and subretinal copic.     Opic dists.     Opic dists.       Non recordable     Cons: 51/2 out of 120UV b wave. Root of 220UV b wave. Root of 220UV b wave. Root of 220UV b wave. Root of and etema     N/A     N	ц	~	z	Y (severe R/G; severe B/Y)	Very small peripheral crystals	Mild lenticular sclerosis	Extreme outer retinal ar. choriocapillaris atrophy small white inner retin crystals		outer retinal and choriocapillaris ny with fine white retinal crystals r in midperiphery, spicular inner I pigment migration with some larger ent patches, small scalloped patches of ved retina in periphery
Y     N     Y (medium R/G     Clear     Clear     Clear     Patchy outer retinal and chicko: sevee B/Y)     Patchy outer retinal and with fine white inner retinal crystals     Patchy outer retinal and white deposits     Patchy outer retinal and choice desci glosis     Patchy outer retinal and choice desci       Y     Y     Y     V     Cear     Contrial opacities     Macular elema and inferior     Optic desci glosis     migration.       Y     Y     Y     V     Contrial opacities     Initially clear, finally     Initially clear, finally     Diffuse crystalline deposis     Copstalline of conscillence     Optic desci       Y     Y     Y     Contra 25% residual function     Optic desci     Optic desci     Optic desci       Conc. 25% residual function     Conc. 25% residual function     Optic desci     Optic desci     Optic desci     Optic desci       Conc. 5.0V out of 12.0V b wave, Rod: 12.0V b wave, Rod: 12.0V b wave, Rod: 12.0V b wave     Contra and subretinal     Optic rocular     Optic rocular     Optic rocular     Optic rocular     Optic rocular     Optic rocular       Non recordable     Conres 5.0V out of 12.0V b wave, Rod: 12.0V b wave, Rod: 12.0V b wave, Rod: 12.0V b wave     Contra and subretinal crows so rocular     Conservation     Conservation     Conservation     Conservation     Conservation     Conservation     Conservation     Conservation     Conservation <td>IJ</td> <td>~</td> <td>Y (mild)</td> <td>N/A</td> <td>Clear</td> <td>Pseudophakic</td> <td>Severe atrophy, crystals yellow white flecks</td> <td>Δ_</td> <td>itary changes</td>	IJ	~	Y (mild)	N/A	Clear	Pseudophakic	Severe atrophy, crystals yellow white flecks	Δ_	itary changes
Y       Y       Y       Clear       Contral opacties       Macular edema and inferior       Optic disks optic disks optic disks optic disks optic disks option         Y       Y       Y       Y       Y       Optic disks optic disks option       Optic disks option         Y       Y       Y       Y       Optic disks optic disks option       Optic disks option         Feature       Electrophysiology       Crystalline deposits       Initially clear, finally       Optic disks optic disks option       Optic disks optic disks option         Electrophysiology       Cone: 25% residual function, root: 20% residual function       OCT: Initiaterihal and subrethial option       Optic disks optic	т	≻	Z	Y (medium R/G defect; severe B/Y)	Clear	Clear	Patchy outer retinal and choriocapillaris atrophy with fine white inner retinal crystals		outer retinal and choriocapillaris atroph ine white retinal crystals mostly in eriphery, spicular inner retinal pigment tion, with some larger pigment patches
Y       Y       Y       Crystaline deposits       Initially clear, finally and reduced refixess       Consumption up, pointanity subcapsular       Infuse crystaline deposits       Follow up, pointanity subcapsular       Infuse crystaline deposits       Constaline deposits <td>_</td> <td>~</td> <td>~</td> <td>~</td> <td>Clear</td> <td>Cortical opacities</td> <td>Macular edema and infe gliosis</td> <td>0</td> <td>lisks vital, narrow vessels, RPE atrophy, lline deposits</td>	_	~	~	~	Clear	Cortical opacities	Macular edema and infe gliosis	0	lisks vital, narrow vessels, RPE atrophy, lline deposits
Electrophysiology         OCT/FAF         Other ocular         C           Cone: 25% residual function, rod: 20% residual function         Other ocular         O         NA         P           Cone: 50% residual function         OCT.intraretinal and subretinal         NA         P         P           Cone: 5uV out of 120uV b wave, Rod: 12uV out of 220uV b wave         CotT.intraretinal and subretinal         NA         P           Non recordable         Non recordable         NA         NA         P           Non recordable         OD: huge macular hole and nutraretinal and subretinal crystals         NA         P           Non recordable         Non recordable         NA         P         P           Somewhat delayed Tr; infigred exotopic and photopic responses on the lower normal limits, critical atrophy, rostalline deposits         P         P <t< td=""><td>_</td><td>~</td><td>~</td><td>~</td><td>Crystalline deposits</td><td>Initially clear, finally primarily subcapsular post cataract</td><td>Diffuse crystalline depos and reduced reflexes</td><td>-</td><td>line deposits, progressive atrophy durin r up, finally choroideremia-like fundus</td></t<>	_	~	~	~	Crystalline deposits	Initially clear, finally primarily subcapsular post cataract	Diffuse crystalline depos and reduced reflexes	-	line deposits, progressive atrophy durin r up, finally choroideremia-like fundus
Come: 55% residual function, rod: 20% residual function     FAF: Patchy hypofluorescence     N/A     P       rod: 20% residual function     CCT: intraretinal and subretinal     N/A     P       Come: 5uV out of 120uV b wave, Rod: 12uV out of 220uV b wave     FAF: Patchy hypofluorescence     N/A     P       Non recordable     Non recordable     N/A     P       Non recordable     OCT: intraretinal and subretinal crystals     N/A     P       Non recordable     N/A     P     P       Non recordable     OD: huge macular hole and mole and and anothy. crystalline     P       Semewhat delayed IT, in outer rings IT delayed, amplitude     Cuter retinal atrophy with fine     N/A     P       Subnormal     ERG: absent rod-specific responses     <	Family	Electrophysi	iology			OCT/FAF	Other	ocular	Other systemic
Cone: SuV out of 120V b wave, Rod: 12uV out of 220uV b wave       Foveal thinning, remodeling, mMA       N         Non recordable       Non recordable       N/A       P         Non recordable       OD: huge macular hole and subretinal crystals       N/A       P         Non recordable       OD: huge macular hole and nuck       N/A       P         Non recordable       OD: huge macular hole and nuck       N/A       P         Stational       OD: huge macular hole and nuck       N/A       P         OD: huge macular hole and nuck       OD: huge macular hole and nuck       N/A       P         Stanzfeld: scotopic and photopic responses on the lower normal imits, crystalline deposits       Patient notes       P         Somewhat delayed IT, mfERG: central responses with almost deposits, central retina (fovea)       central vision defects       and difficultyreading         Normal       ERG: absent rod-specific responses       Outer retinal atrophy with fine       N/A       N         ERG: absent rod-specific and cone-specific responses       Conter retinal atrophy with fine       N/A       N         ERG: absent rod-specific and cone-specific responses       Cuter retinal atrophy with fine       N/A       N         ERG: absent rod-specific and cone-specific responses       Cuter retinal atrophy with fine       N/A       N         Reti	<	Cone: 25% rod: 20% r	residual function residual function			FAF: Patchy hypofluorescen OCT: intraretinal and subre crystals and edema	la		N/A
Non recordable       Non recordable       N/A       N/A         Non recordable       OD: huge macular hole and       N/A       N/A         Non recordable       OD: huge macular hole and       N/A       N/A       N/A         Of anzfeld: scotopic and photopic responses on the lower normal limits, somewhat delayed IT, in outer rings IT delayed, amplitude       Retinal atrophy, crystalline       Patient notes       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       Patient notes       N/A       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A       P       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A       P         RG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A       P         RG: absent rod-specific networes       Outer retinal atrophy with fine       N/A       P         retinal rot-specific responses       Outer retinal atrophy with fine       N/A       P	в	Cone: 5uV	out of 120uV b v	vave, Rod: 12uV out o	ıf 220uV b wave	Foveal thinning, remodeling intraretinal and subretinal (	rvstals		N/A
Ganzfeld: scotopic and photopic responses on the lower normal limits, somewhat delayed IT; mfERG: central responses with almost mormal amplitudes and IT, in outer rings IT delayed, amplitude       Retinal atrophy, crystaline       Patient notes         somewhat delayed IT; mfERG: central responses with almost intact photoreceptors       and difficultyreading         normal amplitudes and IT, in outer rings IT delayed, amplitude       with almost intact photoreceptors       and difficultyreading         RGG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A         ERG: absent rod-specific and cone-specific responses       Outer retinal atrophy with fine       N/A         retinal crystals at all levels, crystals most visible on infrared and redfree reflectance imaging, blue light fundus autofluorescence wirtually absent       N/A	υD	Non record: Non record:	able able			Retinal atrophy, crystalline c OD: huge macular hole and neurosensoric detachment OS: retinal atrophy, crystalline deposits	eposits		N/A Depression, elevated blood pressure, elevated cholesterol levels
ERG: absent rod-specific and cone-specific responses Outer retinal atrophy with fine N/A retinal crystals at all levels, crystals most visible on infrared and redfree reflectance imaging, blue light fundus autofluorescence virtually absent	ш	Ganzfeld: s somewhat normal am subnormal	cotopic and phot delayed IT; mfER plitudes and IT, ii	opic responses on the l (G: central responses w n outer rings IT delayec	lower normal limits, /ith almost d, amplitude	Retinal atrophy, crystalline deposits, central retina (fov with almost intact photore	F	notes al vision defect: ifficultyreading	
	ц	ERG: absent	t rod-specific and	l cone-specific response	81	Outer retinal atrophy with f retinal crystals at all levels, crystals most visible on infr and redfree reflectance im: blue light fundus autofluor virtually absent	d ng, ence		N/A

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21

amily	Family Electrophysiology	OCT/FAF	Other ocular	Other systemic
σ	Full-field ERG: Dark adapted ERG 55% of normal, light adapted 45% of normal, flicker 40% of normal; mfERG: severe central amolitude reduction	OCT not available, FAF multiple patchy areas of intensity loss	N/A	WA
т	ERG: absent rod-specific responses; residual cone-specific responses	Outer retinal atrophy with fine retinal crystals at all levels, crystals most visible on infrared and redfree reflectance imaging, blue light fundus autofluorescence shows hypofluorescent scalloped patches surrounded by thin interconnecting bridges of normo- and hyperautofluorescence	MA	MA
	Ganzfeld ERG extinguished, mfERG central emaining responses	Cystoid macular edema, crystalline deposits	N/A	Hearing problems, thalassemia minor, chronic nephropathy
	Initially negative scotopic ERG. Finally nondetectable.	Deposits in deep retinal layer, obscured in fluorescein angiography	N/A	N/A



**Figure 3.** Crystalline deposits seen on the lens (i) and corneal limbus (ii) of 54-year-old proband D with compound heterozygous mutations in *CYP4V2* c.802-8\_810delinsGC; p.(?), p.(Arg400Cys).

pathogenic by the two in silico programs SIFT and Poly-Phen (Table 2). The p.(Thr417Asnfs\*2) frameshift mutation is presumed to create a premature stop codon one amino acid residue downstream, and in addition might be targeted for nonsense-mediated decay (NMD). In silico prediction of the c.604+4A>G splice site mutation showed a decrease in the strength of the splice donor site due to the alteration (e.g., 12% decrease in SpliceSite finder-like, 52% in MaxEntScan, 89.5% in NNSPLICE, and 9% decrease in Human Splicing Finder [HSF]), suggesting that this mutation might alter *CYP4V2* splicing.

Upon assessing the segregation of mutations in available family members, an inconsistency was noted for proband F, who carried an apparent homozygous mutation c.1249dup; p.(Thr417Asnfs\*2). The mutation was found to segregate in the patient's mother however not in her father, while paternity was confirmed. Subsequently, genomic qPCR analysis revealed a heterozygous deletion encompassing all exons of CYP4V2, in both proband F and her father (Fig. 2A). To further delineate the breakpoints and determine the extent of this new deletion, SNP-chip analysis was performed. The deletion size was demonstrated to vary between 3.8 and 4.1 Mb, spanning the entire CYP4V2 gene and several other genes, including two OMIM genes (KLKB1 and F11) (Fig. 2B). No CYP4V2 deletions were detected in the two other probands who carried a single heterozygous allele (data not shown).

#### **Patient phenotypes**

The clinical data for all 10 patients that carried one or two *CYP4V2* alleles are described in Table 3. Disease onset in these patients ranged from 18 to 57 years of age, with duration of illness ranging from 8 to 33 years. Visual acuities (VA) varied extensively and ranged from 20/20 to light perception (LP), with large discrepancies in VA between both eyes occasionally noted. With the exception of three patients with no refractive error, all subjects were myopic with astigmatism with refraction ranging from -0.50 to -6.00 dioptres. Goldmann visual fields also revealed a wide spectrum of severity, ranging from relatively normal to severe constriction (5–10° remaining). Central, paracentral, and ring scotomas were also noted. Crystalline deposits were noted in the corneal limbus of three patients. In proband D, crystalline deposits were seen on the lens as well as the limbus (Fig. 3). Posterior subcapsular cataracts (PSCC) and cortical opacities were noted in several patients.

The predominant findings on fundus examination were extensive choroidal sclerosis, crystalline intraretinal deposits, attenuated vessels, and intraretinal spicular pigmentation in varying degrees of severity (Fig. 4). Macular atrophy was often seen. Of note is proband A, whose fundus examination revealed choroidal sclerosis and peripheral pigmentation but no crystal deposits. On OCT, however, crystal deposits were observed in the subretinal space (Figs. 4A and 5A). Proband J was similarly reported to have a "choroideremia-like" fundus appearance, with deposits seen in the subretinal layer on OCT. Proband G was also noted to have an unusual fundus appearance, consisting of large discoid patches of atrophy separated by ridges of relatively intact tissue, along with the presence of choroidal sclerosis and crystalline deposits (Fig. 4G).

On OCT, two types of crystals were visualized in patients – tiny intra- and inner retinal crystals and larger, encapsulated subretinal crystals. All nine patients who had OCT were noted to have fine inner retinal crystals; five among these patients also showed larger subretinal crystals (Fig. 5A, B, D, and E, no image available for proband J). Foveal thinning and remodeling of retinal architecture were other pertinent findings. In addition, cystoid macular edema (CME) was documented in two patients. ERG was abnormal in all patients who underwent testing, ranging from nonrecordable to subnormal photoreceptor function.

Disease progression was noted in most patients for whom previous clinical data were available, with declining VA and visual fields. Progression of disease in the fundus of proband F over 18 years was documented photographically (Fig. 4F) and shows the development of the choroidal sclerosis and the clinical disappearance of the retinal crystals.

The clinical presentation of patients without identified *CYP4V2* mutations was indistinguishable from that of the *CYP4V2*-positive patients. The phenotypes of five of the *CYP4V2*-negative patients are shown in Figures S1 and S2. These patients demonstrated the same severe changes that were present in the patients with *CYP4V2* mutations, including choroidal show and sclerosis, the diffuse nature of disease, bone spicules, and macular involvement with or without crystalline deposits. OCT images similarly

reveal the fine inner retinal crystals as well as the larger sometimes encapsulated subretinal crystals.

## Discussion

We identified eight different CYP4V2 mutations in 10/19 patients in our cohort of crystalline retinal dystrophies, clinically diagnosed as Bietti crystalline dystrophy. We identified four novel mutations and one genomic rearrangement (deletion). This is the first large deletion reported for BCD. Eight patients were found to have two mutations in CYP4V2, while in two patients only one mutation was detected, despite extensive analysis of the coding region using Sanger sequencing and CNV analysis. Until now, 57 mutations in CYP4V2 have been associated with BCD. These include 43 missense/nonsense mutations, eight splicing mutations, four small deletions, one small insertion, and one small indel (Li et al. 2004; Lee et al. 2005; Lin et al. 2005; Shan et al. 2005; Wada et al. 2005; Jin et al. 2006; Lai et al. 2007; Zenteno et al. 2008; Mamatha et al. 2011; Xiao et al. 2011; Yokoi et al. 2011; Haddad et al. 2012; Manzouri et al. 2012; Parravano et al. 2012; Song et al. 2013; Halford et al. 2014; Yin et al. 2014). To our knowledge, we are the first to identify a deletion including CYP4V2 and several other genes in a patient with BCD, expanding the molecular pathogenesis of BCD. Interestingly, previous studies have reported five patients in which only single heterozygous mutations were found in CYP4V2, possibly implicating CYP4V2 CNVs as found in proband F (Li et al. 2004; Shan et al. 2005; Jin et al. 2006; Rossi et al. 2013).

The pathogenic role of CNVs and the importance of their detection has been previously described in recessive retinal dystrophies such as retinitis pigmentosa, Leber congenital amaurosis (LCA), and cone dystrophy with supernormal rod response (CDSRR) (Wissinger et al. 2011; Eisenberger et al. 2013). Ultimately, the detection of such CNVs clarifies diagnosis, and may influence genetic counseling. Our results demonstrate that CNV is a mutational mechanism that can lead to BCD as well, and that screening for such variations may be necessary to avoid diagnostic uncertainties.

The deletion in proband F also covers two genes *KLKB1* and *F11*, both of which are involved in the contact activated coagulation pathway. *KLKB1* encodes plasma prekallikrein – also known as Fletcher factor – a glycoprotein in the kallikrein–kininogen–kinin system (KKS) which participates in coagulation, fibrinolysis, and inflammation. Plasma prekallikrein is converted to the serine protease plasma kallikrein by factor XIIa through cleavage of an internal Arg-Ile bond (Chung et al. 1986). Kallikrein then cleaves both low- and high-molecular-weight kininogens to release bradykinin and lys-bradyki-

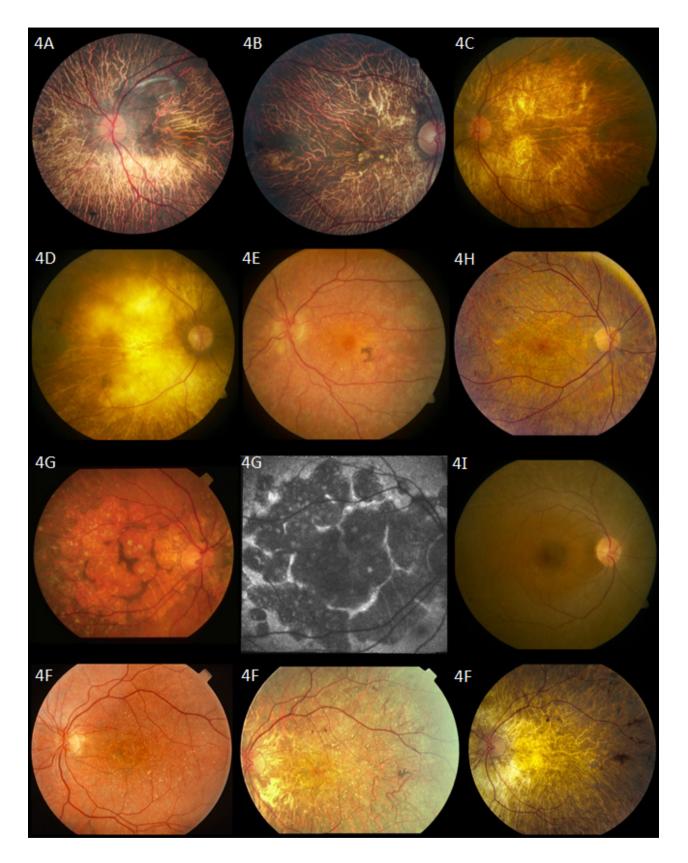
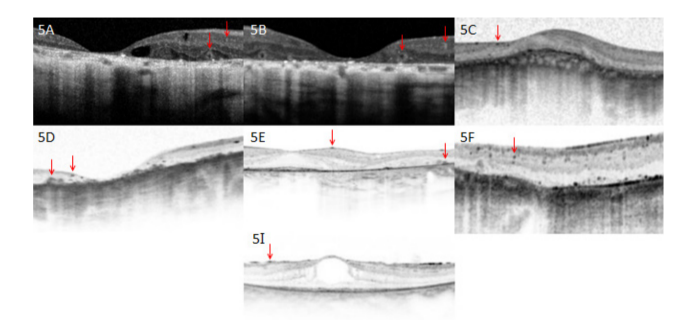


Figure 4. Fundus photographs. Proband A: 57-year-old patient with homozygous mutations in CYP4V2 p.(Ile111Thr). Fundus photograph shows choroidal sclerosis and peripheral pigmentation. No crystal deposits are seen in this patient's fundus. Proband B: 67-year-old patient with homozygous mutations in CYP4V2 p.(Glu202Lys). Fundus photograph shows severe choroidal sclerosis involving the entire macula and fovea, narrowing of the retinal vasculature, with relatively normal optic nerve appearance. Several crystalline deposits can be seen in the macula. Proband C: 47-year-old patient with homozygous mutation in CYP4V2 c.604+4A>G; p.(?). Severe choroidal sclerosis and crystalline deposits in macula. Proband D: 54-year-old patient with compound heterozygous mutations in CYP4V2 c.802-8\_810delinsGC; p.(?), p.(Arq400Cys). Severe choroidal sclerosis and area of visible sclera. Few crystalline deposits seen in midperiphery. Proband E: A 41-year-old patient with compound heterozygous mutations in CYP4V2 p.(Thr81Arg), p.(Arq400Cys). Mild choroidal sclerosis, crystalline deposits, and pigment clump. Some maculopathy is seen. Retinal vasculature remains relatively normal. Proband F: 49-year-old patient with homozygous mutations in CYP4V2 p.(Thr417Nfs\*2). Fundus photographs show progression of disease in left eye (left to right). Of note is the progressive choroidal sclerosis, vascular attenuation and decreasing presence of crystalline deposits over time. Photos taken at ages 31, 38, and 49, respectively. Proband G: 77-year-old patient with compound heterozygous mutation in CYP4V2 c.802-8\_810delinsGC; p.(?), p.(Arq400Cys). Unusual fundus appearance (left) showing large, similarly-sized discoid patches of atrophy separated by small ridges of intact tissue with presence of crystals and choroidal sclerosis. Optic disk and retinal vasculature appear relatively normal. Fundus autofluorescence (middle) confirms areas of atrophy with patches of decreased autofluorescence surrounded by walls of lipofuscin metabolism. Proband H: 38-year-old patient with homozygous mutations in CYP4V2 p.(Arq465Gly). Choroidal sclerosis with diffuse crystalline deposits and pigment clumps in midperiphery. Proband I: 33-year-old patient with one heterozygous mutation in CYP4V2 p.(lle111Thr). Fundus photograph reveals nondescript retina with few crystalline deposits.

nin, mediating the effects of the KKS (Sainz et al. 2007). Recessively inherited mutations in *KLKB1* are known to cause plasma prekallikrein deficiency, resulting in a prolonged activated partial thromboplastin time (aPTT) without increased bleeding tendency in affected patients (Lombardi et al. 2003). Heterozygous patients have been reported to have decreased prekallikrein activity but normal aPTT values (Wynne Jones et al. 2004).



**Figure 5.** Optical coherence tomography images. Proband A: 57-year-old patient with homozygous mutations in *CYP4V2* p.(Ile111Thr). Optical coherence tomography shows severe foveal thinning and cystoid macular edema. Of note are two types of crystal that can be seen; tiny intraretinal crystals and large encapsulated subretinal crystals (red arrows). Proband B: 67-year-old patient with homozygous mutations in *CYP4V2* p.(Glu202Lys). Shown are remarkable thinning of the fovea and remodeling of retina, as well as presence of both intraretinal and subretinal crystals. Proband C: 47-year-old patient with homozygous mutation in *CYP4V2* c.604+4A>G; p.(?). Remodeling of retina with almost complete loss of architecture is seen, as well as tiny intraretinal crystals. Proband D: 54-year-old patient with compound heterozygous mutations in *CYP4V2* c.802-8\_810delinsGC; p.(?), p.(Arg400Cys). Optical coherence tomography shows severe foveal thinning, remodeling of the retina, as well as both intraretinal crystals and larger subretinal crystals. Proband E: A 41-year-old patient with compound heterozygous mutations in *CYP4V2* p.(Thr81Arg), p.(Arg400Cys). Presence of both intraretinal and subretinal crystals. Photoreceptors in fovea remain intact. Proband F: 49-year-old patient with homozygous mutations in *CYP4V2* p.(Thr417Nfs\*2). Optical coherence tomography shows fine intraretinal crystals. Proband I: 33-year-old patient with one heterozygous mutation in *CYP4V2* p.(Ile111Thr). Optical coherence tomography reveals presence of intraretinal crystals.

Factor XI is a homodimeric glycoprotein that circulates in plasma as a noncovalent complex with high-molecularweight kininogen (Fujikawa et al. 1986). Factor XI is activated by factor XII, α-thrombin, and factor XI (autoactivation); it participates in contact activated coagulation by catalyzing the conversion of factor IX to factor IXa and thereby sustaining thrombin generation (Naito and Fujikawa 1991). Factor XI deficiency (sometimes referred to as "hemophilia C") is a rare mild-to-moderate bleeding disorder associated with mutations in F11. Both autosomal recessive and dominant modes of inheritance have been described, the latter possibly being secondary to dominantnegative mutations resulting in mutant proteins forming nonsecretable heterodimers with wild-type subunits (Kravtsov et al. 2004). Moreover, earlier studies demonstrated considerable variability in rates of bleeding in heterozygotes, with some studies even noting no distinction between homozygotes and heterozygotes (Ragni et al. 1985; Bolton-Maggs et al. 1988). Upon revisiting proband F's medical records, we found no history of coagulopathy.

In addition to the two patients in whom only single mutations were detected, we failed to identify any CYP4V2 mutations in nine other patients. The rate of CYP4V2 mutation detection demonstrated in our study was relatively low, especially when compared to a previous study by Xiao et al. (2011), in which CYP4V2 mutations were found in >95% of BCD patient families (Xiao et al. 2011). Our results suggest that CYP4V2 mutations in our patients may reside in genomic locations that were not studied in our gene sequencing protocol, that is, in promoter, untranslated region, or deep intronic regions. Alternatively our data suggests locus heterogeneity and a second Bietti's gene. We were able to re-examine five CYP4V2-negative crystalline retinal dystrophy patients and found that their phenotypes (Figs. S1, S2) are indistinguishable from our BCD patients.

In accordance with previous studies, we found considerable phenotypic variability in our cohort, with no obvious correlation of severity or features with patient demographics. Although disease progression was noted in most patients (for whom historical clinical data were available), length of progression was not clearly associated with a more severe phenotype. Previously, it has been postulated that environmental factors affecting lipid metabolism (e.g., diet) may also contribute to the spectrum of disease (Lee et al. 2005; Rossi et al. 2013). Significant clinical variability was also noted for patients sharing common genotypes, and even within one family. However, it can be observed that in general within our cohort, those with genotypes resulting in splicing defects or nonsense mutations (i.e., c.802-8\_810delinsGC; p.(?), c.604+4A>G; p.(?), and c.1249dup; p.(Thr417Asnfs\*2)) demonstrated greater clinical severity. Notably, proband F

with the genomic deletion was documented to have one of the most clinically severe phenotypes in our cohort. It is possible that the heterozygous *CYP4V2* deletion contributes to her advanced disease state, however this remains unclear as the patient was also found to have a severe frameshift mutation.

Furthermore, proband E, with a mild phenotype despite 21 years of disease, was found to have compound heterozygous mutations, with one allele c.242C>G; (p.Thr81Arg) predicted to be comparatively benign (Table 2).

It should be noted, however, that these correlations between mutation and disease severity have not been consistently shown in previous studies; for instance, although Lai et al. (2007) and Halford et al. (2014) document a similar pattern, Rossi et al. (2013) note that this correlation was not present in their patients. As well, two patients (one with mutation c.1526C>T; (p.P509L) and the other c.1393A>G; (p. p.R465G)) in whom only single mutations were identified in previous studies failed to show the same severity of disease as seen in proband F (Jin et al. 2006; Rossi et al. 2013). The clinical features of the three other heterozygous patients in the literature were not described (Li et al. 2004; Shan et al. 2005).

In our patients, we documented four locations and types of the crystals: small, widespread inner retinal crystals; large, encapsulated subretinal crystals; corneal limbus crystals; and lenticular crystals. The presence of encapsulated subretinal crystals (in addition to inner retinal crystals) on OCT has previously been reported (Pennesi and Weleber 2013). Initially described by Zweifel et al. (2009) in a series of patients with various retinal disorders, these structures were termed "outer retinal tubulation." However, a more recent study with BCD patients found that the hyperrefractive structures identified were in fact spherical (Kojima et al. 2012).

It is of note that proband D with compound heterozygous mutations c.802-8\_810delinsGC; p.(?) and c.1198C >T; p.(Arg400Cys) was found to have crystalline deposits on the lens, which is atypical for BCD. Two separate reports of lenticular crystals have been documented previously, with both patients described carrying the c.802-8\_810delinsGC mutation as well, though in a homozygous state (Yokoi et al. 2010; Chung et al. 2013).

The phenotype of BCD is progressive and severe. In the later stages, as retinal crystals become increasingly difficult to detect on clinical examination and disappear with disease progression, the phenotype can overlap with choroidal sclerosis and some phases of choroideremia; for instance, as reported in the late-stage fundus appearance of probands A and J (Mataftsi et al. 2004; Mansour et al. 2007; Xiao et al. 2011). It is therefore likely that when patients are seen in the later stages of disease, a diagnosis of a nonspecific retinal dystrophy or atypical retinitis pigmentosa may be made. However, in both these patients we documented that OCT reveals nevertheless the presence of retinal crystals. Our results therefore indicate that OCT plays an essential role in the diagnosis and differentiation of BCD from other retinal pathology presenting primarily with choroidal sclerosis.

In conclusion, we identified four novel *CYP4V2* mutations in 10/19 BCD patients as well as the first genomic rearrangement (large deletion) implicating *CYP4V2*. Our results emphasize the importance of CNV screening in BCD. Although the phenotype of the patient with the *CYP4V2* deletion was severe, no clear genotype–phenotype correlation could be established. It would be interesting to note what patterns emerge in future when more patients with large deletions as such are reported. Finally, the nine patients in whom no *CYP4V2* mutations were found suggest that *CYP4V2* mutations may reside in genomic locations that were not covered by our protocol. Alternatively, locus heterogeneity might underlie BCD; further investigations will follow to explore these possibilities.

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## **Conflict of Interest**

None declared.

#### References

Bietti, G. 1937. Ueber familiaeres vorkommen von "retinitis punctata albescens" (verbunden mit "dystrophia marginalis

cristallinea corneae"), glitzern des glaskoerpers und anderen degenerativen augenveraenderungen. Klin. Monbl. Augenheilkd. 99:21.

- Bolton-Maggs, P., B. Wan-Yin, A. McGraw, J. Slack, and P. Kernoff. 1988. Inheritance and bleeding in factor XI deficiency. Br. J. Haematol. 69:521–528.
- Chung, D. W., K. Fujikawa, B. A. McMullen, and E. W. Davie. 1986. Human plasma prekallikrein, a zymogen to a serine protease that contains four tandem repeats. Biochemistry 25:2410–2417.
- Chung, J. K., J. H. Shin, B. R. Jeon, C. S. Ki, and T. K. Park. 2013. Optical coherence tomographic findings of crystal deposits in the lens and cornea in Bietti crystalline corneoretinopathy associated with mutation in the CYP4V2 gene. Jpn. J. Ophthalmol. 57:447–450.
- Collin, R. W., L. I. van den Born, B. J. Klevering, M. de Castro-Miró, K. W. Littink, K. Arimadyo, et al. 2011.
  High-resolution homozygosity mapping is a powerful tool to detect novel mutations causative of autosomal recessive RP in the Dutch population. Invest. Ophthalmol. 52:2227–2239.
- Desmet, F. O., D. Hamroun, M. Lalande, G. Collod-Béroud, M. Claustres, and C. Béroud. 2009. Human Splicing Finder: an online bioinformatics tool to predict splicing signals. Nucleic Acids Res. 37:e67.
- Eisenberger, T., C. Neuhaus, A. O. Khan, C. Decker, M. N. Preising, C. Friedburg, et al. 2013. Increasing the yield in targeted next-generation sequencing by implicating CNV analysis, non-coding exons and the overall variant load: the example of retinal dystrophies. PLoS One 8:e78496.
- Fujikawa, K., D. W. Chung, L. E. Hendrickson, and E. W. Davie. 1986. Amino acid sequence of human factor XI, a blood coagulation factor with four tandem repeats that are highly homologous with plasma prekallikrein. Biochemistry 25:2417–2424.
- Haddad, N. M., N. Waked, R. Bejjani, Z. Khoueir, E. Chouery, S. Corbani, et al. 2012. Clinical and molecular findings in three Lebanese families with Bietti crystalline dystrophy: report on a novel mutation. Mol. Vis. 18:1182–1188.
- Halford, S., G. Liew, D. S. Mackay, P. I. Sergouniotis, R. Holt, S. Broadgate, et al. 2014. Detailed phenotypic and genotypic characterization of Bietti crystalline dystrophy. Ophthalmology 121:1174–1184.
- Hu, D. N. 1987. Prevalence and mode of inheritance of major genetic eye diseases in China. J. Med. Genet. 24:584–588.
- Jin, Z. B., S. Ito, Y. Saito, Y. Inoue, Y. Yanagi, and N. Nao-i. 2006. Clinical and molecular findings in three Japanese patients with crystalline retinopathy. Jpn. J. Ophthalmol. 50:426–431.
- Kojima, H., A. Otani, K. Ogino, S. Nakagawa, Y. Makiyama, M. Kurimoto, et al. 2012. Outer retinal circular structures in patients with Bietti crystalline retinopathy. Br. J. Ophthalmol. 96:390–393.
- Kravtsov, D., W. Wu, C. M. Meijers, M. F. Sun, M. A. Blinder, T. P. Dang, et al. 2004. Dominant factor XI

deficiency caused by mutations in the factor XI catalytic domain. Blood. 104:128–134.

Lai, T. Y., T. K. Ng, P. O. Tam, G. H. Yam, J. W. Ngai, W. M. Chan, et al. 2007. Genotype phenotype analysis of Bietti's crystalline dystrophy in patients with CYP4V2 mutations. Invest. Ophthalmol. Vis. Sci. 48:5212–5220.

Lai, T. Y., K. O. Chu, K. P. Chan, T. K. Ng, G. H. Yam, D. S. Lam, et al. 2010. Alterations in serum fatty acid concentrations and desaturase activities in Bietti crystalline dystrophy unaffected by CYP4V2 genotypes. Invest. Ophthalmol. Vis. Sci. 51:1092–1097.

Lee, J., X. Jiao, J. F. Hejtmancik, M. Kaiser-Kupfer, W. A. Gahl, T. C. Markello, et al. 2001. The metabolism of fatty acids in human Bietti crystalline dystrophy. Invest. Ophthalmol. Vis. Sci. 42:1707–1714.

Lee, K. Y., A. H. Koh, T. Aung, V. H. Yong, K. Yeung, C. L. Ang, et al. 2005. Characterization of Bietti crystalline dystrophy patients with CYP4V2 mutations. Invest. Ophthalmol. Vis. Sci. 46:3812–3816.

Li, A., X. Jiao, F. L. Munier, D. F. Schorderet, W. Yao, F. Iwata, et al. 2004. Bietti crystalline corneoretinal dystrophy is caused by mutations in the novel gene CYP4V2. Am. J. Hum. Genet. 74:817–826.

Lin, J., K. M. Nishiguchi, M. Nakamura, T. P. Dryja, E. L. Berson, and Y. Miyake. 2005. Recessive mutations in the CYP4V2 gene in East Asian and Middle Eastern patients with Bietti crystalline corneoretinal dystrophy. J. Med. Genet. 42:e38.

Lombardi, A. M., M. T. Sartori, L. Cabrio, M. Fadin, E. Zanon, and A. Girolami. 2003. Severe prekallikrein (Fletcher factor) deficiency due to a compound heterozygosis (383Trp stop codon and cys529Tyr). Thromb. Haemost. 90:1040– 1045.

Mamatha, G., V. Umashankar, N. Kasinathan, T. Krishnan, R. Sathyabaarathi, T. Karthiyayini, et al. 2011. Molecular screening of the CYP4V2 gene in Bietti crystalline dystrophy that is associated with choroidal neovascularization. Mol. Vis. 17:1970–1977.

Mansour, A. M., S. H. Uwaydat, and C. C. Chan. 2007. Long-term follow-up in Bietti crystalline dystrophy. Eur. J. Ophthalmol. 17:680–682.

Manzouri, B., P. I. Sergouniotis, A. G. Robson, A. R. Webster, and A. Moore. 2012. Bietti crystalline retinopathy: report of retinal crystal deposition in male adolescent siblings. Arch. Ophthalmol. 130:1470–1473.

Marmor, M. F., A. B. Fulton, G. E. Holder, Y. Miyake, M. Brigell, and M. Bach. 2009. Standard for clinical electroretinography. Doc. Ophthalmol. 118:69–77.

Mataftsi, A., L. Zografos, E. Millá, M. Secrétan, and F. L. Munier. 2004. Bietti's crystalline corneoretinal dystrophy: a cross-sectional study. Retina 24:416–426.

Miller, S. A., D. D. Dykes, and H. F. Polesky. 1988. A simple salting out procedure for extracting DNA from human nucleated cells. Nucleic Acids Res. 16:1215. Naito, K., and K. Fujikawa. 1991. Activation of human blood coagulation factor XI independent of factor XII: factor XI is activated by thrombin and factor XIa in the presence of negatively charged surfaces. J. Biol. Chem. 266:7353–7358.

Nakano, M., E. J. Kelly, and A. E. Rettie. 2009. Expression and characterization of CYP4V2 as a fatty acid omega-hydroxylase. Drug Metab. Dispos. 37:2119–2122.

Ng, P. C., and S. Henikoff. 2003. SIFT: predicting amino acid changes that affect protein function. Nucleic Acids Res. 31:3812–3814.

Parravano, M., M. Sciamanna, P. Giorno, A. Boninfante, and M. Varano. 2012. Bietti crystalline dystrophy: a morpho-functional evaluation. Doc. Ophthalmol. 124: 73–77.

Pennesi, M. E., and R. G. Weleber. 2013. High-resolution optical coherence tomography shows new aspects of Bietti crystalline retinopathy. Retina 30:531–532.

Ragni, M., D. Sinha, F. Seaman, J. Lewis, J. Spero, and P. Walsh. 1985. Comparison of bleeding tendency, factor XI coagulant activity, and factor XI antigen in 25 factor XI-deficient kindreds. Blood 65:719–724.

Reese, M. G., F. H. Eeckman, D. Kulp, and D. Haussler. 1997. Improved splice site detection in Genie. J. Comput. Biol. 4:311–323.

Rossi, S., F. Testa, A. Li, F. Yaylacioğlu, C. Gesualdo, J. F. Hejtmancik, et al. 2013. Clinical and genetic features in Italian Bietti crystalline dystrophy patients. Br. J. Ophthalmol. 97:174–179.

Sainz, I. M., R. A. Pixley, and R. W. Colman. 2007. Fifty years of research on the plasma kallikrein-kinin system: from protein structure and function to cell biology and in-vivo pathophysiology. Thromb. Haemost. 98:77–83.

Shan, M., B. Dong, X. Zhao, J. Wang, G. Li, Y. Yang, et al. 2005. Novel mutations in the CYP4V2 gene associated with Bietti crystalline corneoretinal dystrophy. Mol. Vis. 11:738– 743.

Song, Y., G. Mo, and G. Yin. 2013. A novel mutation in the CYP4V2 gene in a Chinese patient with Bietti's crystalline dystrophy. Int. Ophthalmol. 33:269–276.

Wada, Y., T. Itabashi, H. Sato, M. Kawamura, A. Tada, and M. Tamai. 2005. Screening for mutations in CYP4V2 gene in Japanese patients with Bietti's crystalline corneoretinal dystrophy. Am. J. Ophthalmol. 139:894–899.

Wilson, D. J., R. G. Weleber, M. L. Klein, R. B. Welch, and W. R. Green. 1989. Bietti's crystalline dystrophy. A clinicopathologic correlative study. Arch. Ophthalmol. 107:213–221.

Wissinger, B., S. Schaich, B. Baumann, M. Bonin, H. Jägle, C. Friedburg, et al. 2011. Large deletions of the KCNV2 gene are common in patients with cone dystrophy with supernormal rod response. Hum. Mutat. 32:1398–1406.

Wynne Jones, D., G. Russell, S. L. Allford, K. Burdon, G. A. Hawkins, D. W. Bowden, et al. 2004. Severe prekallikrein deficiency associated with homozygosity for

an Arg94Stop nonsense mutation. Br. J. Haematol. 127:220-223.

Xiao, X., G. Mai, S. Li, X. Guo, and Q. Zhang. 2011. Identification of CYP4V2 mutation in 21 families and overview of mutation spectrum in Bietti crystalline corneoretinal dystrophy. Biochem. Biophys. Res. Commun. 409:181–186.

Yeo, G., and C. B. Burge. 2004. Maximum entropy modeling of short sequence motifs with applications to RNA splicing signals. J. Comput. Biol. 11:377–394.

- Yin, H., C. Jin, X. Fang, Q. Miao, Y. Zhao, Z. Chen, et al. 2014. Molecular analysis and phenotypic study in 14 Chinese families with Bietti crystalline dystrophy. PLoS One 9:e94960.
- Yokoi, Y., M. Nakazawa, S. Mizukoshi, K. Sato, T. Usui, and K. Takeuchi. 2010. Crystal deposits on the lens capsules in Bietti crystalline corneoretinal dystrophy associated with a mutation in the CYP4V2 gene. Acta Ophthalmol. 88:607– 609.
- Yokoi, Y., K. Sato, H. Aoyagi, Y. Takahashi, M. Yamagami, and M. A. Nakazawa. 2011. Novel compound heterozygous mutation in the CYP4V2 gene in a Japanese patient with Bietti's crystalline corneoretinal dystrophy. Case Rep. Ophthalmol. 2:296–301.
- Zenteno, J. C., R. Ayala-Ramirez, and F. Graue-Wiechers. 2008. Novel CYP4V2 gene mutation in a Mexican patient

with Bietti's crystalline corneoretinal dystrophy. Curr. Eye Res. 33:313–318.

Zweifel, S. A., M. Engelbert, K. Laud, R. Margolis, R. F. Spaide, and K. B. Freund. 2009. Outer retinal tubulation: a novel optical coherence tomography finding. Arch. Ophthalmol. 127:1596–1602.

## **Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Figure S1.** Fundus photographs of *CYP4V2*-negative patients. Shown are fundus photographs of five *CYP4V2*-negative Bietti crystalline dystrophy patients. As in those with identified *CYP4V2* mutations, predominant findings include choroidal sclerosis with or without visible crystalline deposits, pigment deposition, and vascular attenuation.

**Figure S2.** Optical coherence tomography images of *CYP4V2*-negative patients. OCT images of five *CYP4V2*-negative patients, with presence of small hyperreflective intraretinal crystals and larger subretinal crystals (red arrows).

Table S1. CYP4V2 primer sequences.

Table S2. Conditions and primer sequences.