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# Autosomal Dominant Retinal Dystrophies Caused by a Founder Splice Site Mutation, c.828+3A>T, in *PRPH2* and Protein Haplotypes in *trans* as Modifiers

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Citation: Shankar SP, Hughbanks-Wheaton DK, Birch G, et al. Autosomal dominant retinal dystrophies caused by a founder splice site mutation, c.828+3A>T, in *PRPH2* and protein haplotypes in *trans* as modifiers. *Invest Ophtbalmol Vis Sci.* 2016;57:349–359. DOI:10.1167/ iovs.15-16965 **PURPOSE.** We determined the phenotypic variation, disease progression, and potential modifiers of autosomal dominant retinal dystrophies caused by a splice site founder mutation, c.828+3A>T, in the *PRPH2* gene.

**M**ETHODS. A total of 62 individuals (19 families) harboring the *PRPH2* c.828+3A>T mutation, had phenotype analysis by fundus appearance, electrophysiology, and visual fields. The *PRPH2* haplotypes in *trans* were sequenced for potential modifying variants and generalized estimating equations (GEE) used for statistical analysis.

**R**ESULTS. Several distinct phenotypes caused by the *PRPH2* c.828+3A>T mutation were observed and fell into two clinical categories: Group I (N = 44) with mild pattern dystrophies (PD) and Group II (N = 18) with more severe cone-rod dystrophy (CRD), retinitis pigmentosa (RP), and central areolar chorioretinal dystrophy (CACD). The *PRPH2* Gln304-Lys310-Asp338 protein haplotype in *trans* was found in Group I only (29.6% vs. 0%), whereas the Glu304-Lys310-Gly338 haplotype was predominant in Group II (94.4% vs. 70.4%). Generalized estimating equations analysis for PD versus the CRD/CACD/RP phenotypes in individuals over 43 years alone with the *PRPH2* haplotypes in *trans* and age as predictors, adjusted for correlation within families, confirmed a significant effect of haplotype on severity (P = 0.03) with an estimated odds ratio of 7.16 (95% confidence interval [CI] = [2.8, 18.4]).

**CONCLUSIONS.** The *PRPH2* c.828+3A>T mutation results in multiple distinct phenotypes likely modified by protein haplotypes in *trans*; the odds of having the CACD/RP-like phenotype (versus the PD phenotype) are 7.16 times greater with a Glu304-Lys310-Gly338 haplotype in *trans*. Further functional studies of the modifying haplotypes in *trans* and *PRPH2* splice variants may offer therapeutic targets.

Keywords: phenotype, genetic modifiers, retinal dystrophy

Mutations in the *PRPH2* gene (OMIM:179605) are known to cause a wide range of autosomal dominant retinal dystrophies (adRD; phenotype MIM: 613105, 608133, 169150, 608161, 608133, 136880), but the source of this variation is poorly understood. We previously identified a *PRPH2* (NM\_000322.4) splice site mutation, c.828+3A>T, in numerous patients with retinal dystrophies.<sup>1</sup> Haplotype analysis revealed that the mutation derives from a common ancestor and its prevalence is due to a founder effect.<sup>2</sup> Individuals with the c.828+3A>T mutation express a *PRPH2* splice variant that was not found in controls and is consistent with abnormal splicing. Clinical diagnoses in individuals with this mutation range from pattern dystrophies (PD) to retinitis pigmentosa (RP), cone-rod dystrophy (CRD), and/or unspecified macular dystrophy.<sup>2</sup> This phenotypic variation is comparable to other mutations in *PRPH2*, for example, a 3-bp deletion of codon

153, and a P210R missense change<sup>3</sup> that were reported to cause multiple phenotypes within the same nuclear families.<sup>4</sup> Another *PRPH2* mutation, R172W, also exhibits a founder effect but with similar phenotypes in most families,<sup>5,6</sup> although intrafamilial variation also has been described in two families.<sup>7,8</sup>

In this study, we evaluated clinical records, fundus photographs, and visual function data to characterize the phenotype of the *PRPH2* c.828+3A>T splice site mutation. Due to the founder effect, this cohort is particularly well suited for identifying possible causes of variation that are not in *cis* to the splice site mutation.

Peripherin/*PRPH2* is a photoreceptor-specific membrane protein that homodimerizes with itself. Polymorphic protein haplotypes in *trans* were considered as modifiers of dominant diseases because the two alleles are expressed at the same time in the same tissues, translation is concurrent, and protein products may follow identical pathways and interact directly. Therefore, sequence variants in the *PRPH2* gene in *trans* that code for polymorphic protein isoforms (haplotypes),  $Glu_{304}$ - $Lys_{310}$ - $Gly_{338}$  ( $G_{910}$ - $A_{929}$ - $G_{1013}$ ),  $Gln_{304}$ - $Arg_{310}$ - $Asp_{338}$  ( $C_{910}$ - $G_{929}$ - $A_{1013}$ ), and  $Gln_{304}$ - $Lys_{310}$ - $Asp_{338}$  ( $C_{910}$ - $G_{929}$ - $A_{1013}$ ), and  $Gln_{304}$ - $Lys_{310}$ - $Asp_{338}$  ( $C_{910}$ - $G_{929}$ - $A_{1013}$ ), coded by three polymorphic cSNPs, rs390659 c.910G>C (GAG->CAG) p.Glu304Gln, rs425876, c.929A>G (AAG->AGG) p.Lys310Arg, and rs434102 c.1013G>A (GGC->GAC) p.Gly338Asp were considered as potential modifiers of the disease phenotype.<sup>2,9,10</sup>

We also considered 3 other candidate genes as modifiers, ROM1, GUCA1A, and NXNL1. ROM1 is a photoreceptorspecific gene that encodes a rod outer-membrane protein localized to the photoreceptor outer segment disk membranes. It has 35% homology to PRPH2 and interacts with PRPH2-forming heterotetramers.<sup>11</sup> Mutations in this gene alone have not been shown conclusively to cause any retinal disease, although a digenic form of RP with a PRPH2 mutation has been reported.<sup>12</sup> Therefore, ROM1 was considered as a strong candidate modifier gene, particularly in individuals with the retinitis pigmentosa phenotype. The GUCA1A gene encodes a photoreceptor-specific guanylate cyclase-activator protein, a novel Ca(2+) binding protein that stimulates synthesis of cGMP in photoreceptors.<sup>13</sup> Mutations in this gene are known to cause autosomal dominant cone dystrophy, making it a likely modifier gene.<sup>14,15</sup> Another gene, NXNL1, encodes a thioredoxin-like protein 6, which is specifically expressed in photoreceptors and required for cone viability.<sup>16</sup> We considered GUCA1A and NXNL1 as potential modifier genes, especially in those patients exhibiting predominant macular disease and cone loss.

### METHODS

#### Subjects and Clinical Studies

All subjects were ascertained in the United States after obtaining written informed consent and all procedures conformed to the Declaration of Helsinki and were approved by the Institutional Review Board (IRB) board of each respective institution as described previously.<sup>2</sup> We studied 19 families (FAM) harboring the splice site mutation. Families 1 to 10 were ascertained by David Birch, PhD, at the Retina Foundation of the Southwest (RFSW; Dallas, TX, USA); Fam11 by Richard Ruiz, MD, at University of Texas Health Science Center (Houston, TX, USA); Fam12 by Jonathon Sears, MD, at Cole Eve Institute (Cleveland, OH, USA): Fam13, Fam15, and Fam17 to Fam19 by Edwin Stone, MD, PhD, at the University of Iowa (Iowa City, IA, USA); Fam14 at the University of Oregon (Eugene, OR, USA) by Richard Weleber, MD; and Fam16 by Gerald Fishman, MD, at the University of Illinois (Chicago, IL, USA). Screening for sequence variants was performed at the Molecular Ophthalmology Laboratory at the University of Iowa and the Laboratory for Molecular Diagnosis of Inherited Eye Diseases at the University of Texas Health Science Center.

Clinical findings were analyzed to determine the phenotype of the individuals and classify them into defined diagnostic groups. Fundus photographs were available for 62 individuals from 19 families with the splice site mutation. To maintain consistent standards, we used electrophysiology data from RFSW alone for analysis. Electroretinographic (ERG) data were available for 25 individuals, dark adaptometry for 24 individuals, and Humphrey visual fields for 10 individuals.

### **Visual Acuity**

Best-corrected visual acuity was recorded using an Early Treatment Diabetic Retinopathy Study (ETDRS) chart or standard Snellen visual acuity charts.

### **Visual Fields**

Visual fields were tested at RFSW using standard spot size 3, program 30-2 and/or 60-4 SITA Fast on a Humphrey field analyzer (Humphrey Instruments, Inc., San Leandro, CA, USA).

### Electrophysiology

Patients were dilated using tropicamide 1% and phenylephrine 2.5% and then dark-adapted for 45 minutes before testing. Usually only the eye with the poorer visual acuity was tested. The ERG was recorded from the dilated, dark-adapted eye following the International Society for Clinical Electrophysiology of Vision (ISCEV) standards as described previously.<sup>17</sup>

### **Dark Adaptometry**

Patients' eyes were dilated using tropicamide 1% and phenylephrine 2.5% and dark-adapted completely for 45 minutes before testing. Final dark-adapted thresholds were measured on a Goldmann-Weekers adaptometer according to standard protocols.<sup>18–20</sup>

### **Fundus Photographs**

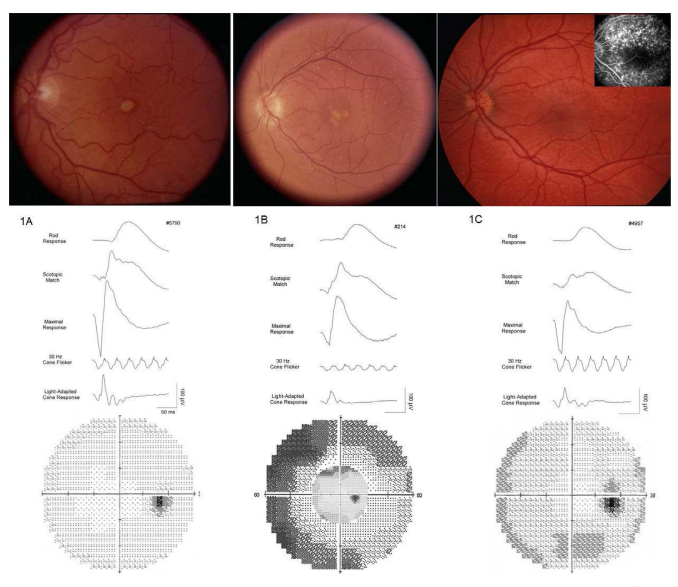
Fundus photographs were obtained from each site whenever available. Phenotypic classification was based on fundus findings available in 62 individuals. All individuals with macular and/or peripheral yellow, orange, or gray (dot and halo) deposits were grouped into various types of PD as described previously.21-23 Patients with macular pigmentation and atrophy preceding peripheral retinal changes were classified as having CRD.<sup>24</sup> If the fundus appeared to have peripheral intraretinal bone spicules, mottling, and granularity of the RPE, and/or had optic nerve pallor and retinal vessel attenuation, patients were classified as having RP.25 Individuals with symmetric, sharply outlined areas of geographic atrophy of the RPE and choroid, approximately 3 to 4 disc diameters in size, and having a yellow-white appearance of the choroidal vessels, were classified as central areolar chorioretinal dystrophy (CACD).26

These diagnoses were combined into two groups to analyze the effect of modifier genes: Group I – milder phenotype with PD: The presence of yellow or pigmented lesions in the retina and/or retinal atrophy confined to the macula alone; and Group II – more severe phenotype with CRD, RP and CACD: These were characterized by photoreceptor loss, RPE, and choroidal atrophy extending outside the macula and/or presence of intraretinal bone spicules.

#### **Molecular Studies**

DNA extraction and screening by SSCP or bidirectional fluorescent di-deoxy sequencing of the genes, *PRPH2* (NM\_000322.4), *ROM1* (NM\_000327.3), *GUCA1A* (NM\_000409.3), and *NXNL1* (NM\_138454.1) were performed using standard conditions as described previously.<sup>1,27</sup> *PRPH2* haplotypes in *trans* were determined by subtraction of the known c.828+3A>T haplotype from the genotype determined via sequencing. Primers used in the study are available in Supplementary Table S1.

#### PRPH2 Retinal Dystrophies



**FIGURE 1.** Fundus photographs, ERG, and Humphrey visual fields of three PD patients. (**A**) Adult-onset foveomacular vitelliform-like PD: 46-year-old female (Fam7-5790) showing lipofuscin deposits in the macula, normal rod and cone responses, very minimal field loss. (**B**) Butterfly-like PD: 49-year-old female (Fam1-214) showing lipofusin deposits in the macula and in the periphery, decreased rod and cone responses, moderate peripheral field loss. (**C**) Flavimaculatus-like PD: 28-year-old female (Fam2-4957) showing peripheral deposits (seen better in fluorescein *inset*), decreased rod responses but normal cone responses, mild peripheral field loss.

#### **Statistical Analysis**

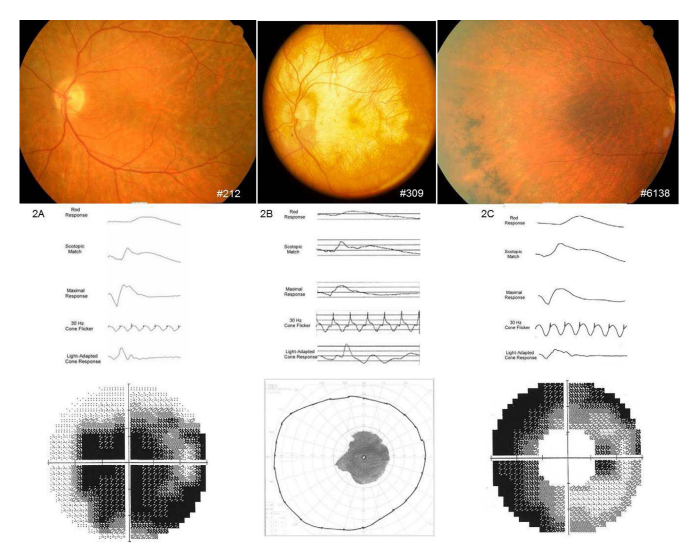
Linear regression was used to analyze the relation between age and visual function parameters in patients with the splice site PRPH2 mutation. The best-fit values of the slope and intercept, along with their standard errors and 95% confidence intervals (CIs), were calculated. Associations between the two phenotype groups and the three protein haplotypes in trans were tested by Fisher's exact probability test. To confirm that significant associations were not due to age differences or familial clustering of genotypes, generalized estimating equation (GEE) models<sup>28</sup> were fit using the "geepack" R package.<sup>29</sup> These regressions modeled phenotype group as a function of haplotype and age via a logistic link, and robustly accounted for genetic correlation within families. An exchangeable correlation structure was specified for each familial cluster, but it is important to note that GEE models are robust even if the correlation structure is misspecified. For all haplotype analyses, we excluded all individuals younger than 43 years,

taking into consideration the fact that the youngest age of an individual with the more severe phenotype in our cohort was 43 years.

#### RESULTS

The presenting symptoms of the individuals with the c.828+3A>T splice site mutation include reduced, altered, or distorted central vision, sensitivity to bright lights, and reduced night vision.

Four distinct phenotypes were observed consistent with PDs, CRD, RP, or CACD with extensive intra- and interfamilial variation. Three types of PD, adult-onset vitelliform macular dystrophy (AVMD), butterfly-shaped macular dystrophy (BMD), and adult-onset flavimaculatus-like macular dystrophy (AOFMD) of three individuals in their 40s are shown in Figures 1A through 1C. Individuals having AVMD and BMD exhibited a milder phenotype at age 40 than the individual with AOFMD,



**FIGURE 2.** Fundus photographs, ERG, and visual fields in more severe retinal dystrophies. (A) Cone-rod dystrophy in a 58-year-old female (Fam1-212) showing photoreceptor and RPE atrophy, reduced rod and cone responses with central field loss, and relative preservation of the peripheral field. (B) Central areolar chorioretinal dystrophy in a 56-year-old male (Fam1-309) showing geographic atrophy of photoreceptors, RPE, and choriocapillaries atrophy, with sparing of peripheral retina with corresponding Goldmann III4e visual field. (C) Retinitis pigmentosa in a 60-year-old female (Fam8-6138) showing pigmentary changes in the periphery with a generalized loss of photoreceptors and RPE, loss of rod function more than cone, and peripheral visual field loss with relative central sparing.

who showed more severe ERG and field loss at age 28 years. A fluorescein angiogram shows hyperfluorescent lesions that are not visible by ophthalmoscope (Fig. 1C, inset). The absence of "dark choroid" distinguishes these from Stargardt's and fundus flavimaculatus caused by mutations in the *ABCA4* gene.<sup>30</sup> Individuals with more extensive vision loss, as in CRD, CACD, and RP phenotypes, are shown in Figures 2A through 2C.

The fundus features of 62 individuals, along with their diagnoses, age, and sex, and the *PRPH2* mini-haplotypes in *trans*, are shown in Table 1 and pedigrees for 14 families are available in the Supplementary Material. Of the individuals, 44 were graded as Group I and had the clinical diagnosis of PD, and 18 were graded as Group II and had CRD, RP, or CACD.

#### Age

The age range of our patients was 16 to 90 years, with majority in the 40- to 50-year range. The youngest individual examined, a 16-year-old male, had minimal yellow lesions in the peripheral retina and the oldest individual, a 90-year-old female, had geographic atrophy in the macula; both belonged to the same family. The majority of the individuals with the splice site mutation fell into the PD Group I (44), with ages ranging from the 20s to the 70s (Table 1). Individuals having a fundus appearance suggestive of CRD (8), RP (6), and CACD (4) were in their fifth to sixth decades of life (Table 1).

#### **Visual Acuity**

Visual acuity data were available for 46 individuals with the splice site mutation, and the best-corrected visual acuity of the better eye was used for plotting (Fig. 3A). Most of the PD patients retain fairly good visual function, with their acuity remaining above 20/40 (0.5) until 65 years of age (green symbols). There was a modest decrease of visual acuity in PD with age due to photoreceptor and RPE atrophy in the macular region (Fig. 3A, solid line). In contrast, patients with CACD and CRD exhibited significantly lower visual acuity (red symbols), with more than half of the patients over age 45 having VA below 20/100 (0.2). Retinitis pigmentosa patients had VA ranging from 20/40 to 20/400 (0.2 to <0.08).

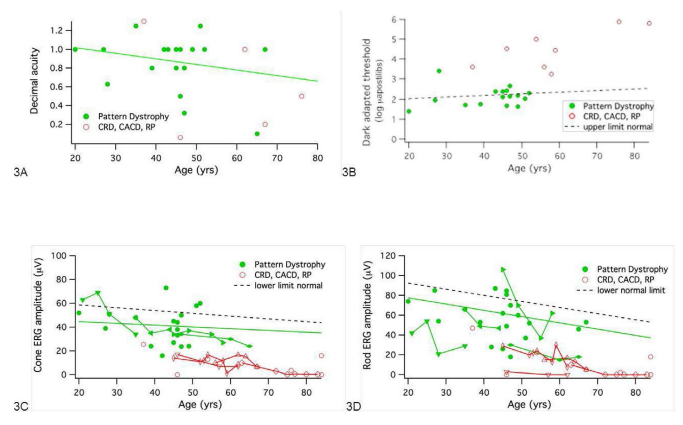
 TABLE 1. Phenotype and Haplotypes of PRPH2 Gene in trans to the Splice Site Mutation

No.	Individual ID	Haplotype in <i>trans</i>	Age, y First Visit	Sex	Phenotype – Clinical Features
1	Fam13-3	C-A-A	16	М	PD - few lipofuscin deposits
2	Fam13-35	G-A-G	24	F	PD - lipofuscin deposits
3	Fam14-7	C-A-A	25	Μ	PD - few lipofuscin deposits
4	Fam13-68	C-A-A	26	F	PD - few lipofuscin deposits
5	Fam5-5738	G-A-G	27	F	PD - lipofuscin deposits in macula and ST arcade
6	Fam2-4957	G-A-G	28	F	PD - granular appearance in macula, flecks in peripapillary region and arcades, confluent, pigment +
7	Fam14-16	G-A-G	30	F	PD – lipofuscin deposits in macula and around arcades
8	Fam2-5878	C-A-A	30	М	PD - minimal lipofuscin deposits
9	Fam8-6226	G-A-G	35	F	PD - grayish yellow lesion in macula with min flecks
10	Fam13-91	G-A-G	37	М	PD – minimal macular lesion
11	Fam13-41	G-A-G	38	F	PD - subtle adult vitelliform lesions in fovea, lipofuscin accumulation near the arcades
12	Fam13-92	G-A-G	38	М	PD – macular pigmentary changes with peripheral drusen-like subretinal deposits
13	Fam14-21	G-A-G	39	М	PD - yellow deposit in macula with min flecks
14	Fam6-5755	G-A-G	39	М	PD - yellow deposit in macula, flecks in peripapillary region and arcades
15	Fam12-3	C-A-A	40	F	PD - yellow deposit in macula, RPE atrophy with flecks
16	Fam7-7082	G-A-G	41	М	PD - yellow deposit in macula with min flecks
17	Fam11-3	G-A-G	41	F	PD - yellow deposits in macula
18	Fam2-9205	G-A-G	43	М	CRD – extensive RPE atrophy in macula, flecks around atrophic zone
19	Fam14-9	G-A-G	45	F	PD - pattern-like maculopathy
20	Fam1-177	C-A-A	45	F	PD - min Flecks, hypopigmentation in macula
21	Fam14-14	G-A-G	46	F	PD – pattern-like maculopathy
22	Fam5-9226	G-A-G	46	F	PD – extensive confluent flecks
23	Fam7-5790	C-A-A	46	F	PD - vitelliform-like lesion at macula, min flecks
24	Fam13-39	G-A-G	47	F	PD - very early pattern lesion in fovea, abnormal lipofuscin deposits beyond the arcades
25	Fam1-262	G-A-G	48	М	RP - extensive atrophy - bone spicules all quadrants
26	Fam12-2	G-A-G	49	М	PD - yellow lesion in macula and flecks
27	Fam14-15	G-A-G	49	F	PD – pattern-like maculopathy
28	Fam1-214	G-A-G	49	F	PD - lipofuscin deposits - macula and periphery
29	Fam1-194	C-A-A	49	F	PD - lipofuscin deposits in macula, min flecks
30	Fam7-6221	C-A-A	49	F	PD - grayish yellow deposits in macula with min flecks
31	Fam8-9048	G-A-G	50	М	PD - multiple flecks, flavimaculatus-like, mottling of RPE
32	Fam2-9207	G-A-G	51	F	PD - yellow lesion at macula, min flecks
33	Fam4-5249	G-A-G	51	F	PD - gray, yellow lesions in macula, min flecks
34	Fam13-6	G-A-G	53	F	CRD - extensive RP and choroidal atrophy in macula and periphery with bone spicules
35	Fam2-4920	G-A-G	54	F	CRD - extensive RP and choroidal atrophy in macula and periphery with bone spicules
36	Fam12-1	G-A-G	55	F	PD - yellow lesions in macula, RPE degeneration with flecks
37	Fam13-18	G-A-G	56	М	PD - macular dot and halo lesion, lipofuscin deposits around arcades
38	Fam5-9234	G-A-G	57	F	CRD - chorioretinal atrophy in macular region min flecks
39	Fam15-1	G-A-G	57	F	RP - initially ad retinal dystrophy with flecks; 2 y later RPE atrophy w/ clumped RPE - pavingstone and photoreceptor degeneration
40	Fam13-1	G-A-G	58	М	PD - fleck-like collections of lipofuscin in macula extending beyond arcade
41	Fam1-302	G-A-G	58	F	PD - yellow flecks in macula
42	Fam1-212	G-A-G	58	F	CRD – extensive atrophy macular region and around peripapillary region
43	Fam13-17	G-A-G	59	М	PD - large reticular lipofuscin deposits extending from the arcades to the perifoveal area
44	Fam14-3	G-A-G	59	М	CACD - central RPE and choroidal atrophy
45	Fam15-6	G-A-G	59	F	PD - lipofuscin deposits, mild arterial attenuation
46	Fam8-6138	G-A-G	60	F	RP - macula - hypopigmentation, peripheral BSs, and drusen-like deposits
47	Fam15-16	G-A-G	61	М	CRD - RPE atrophy w/pigment clumps, arterial attenuation, cobblestone
48	Fam13-34	C-A-A	65	F	PD – 5-pointed butterfly lesion in macula
49	Fam1-365	G-A-G	65	М	PD - yellow-gray lesion with confluent flecks
	Fam13-87	G-A-G	65	F	CACD - multifocal choroidal atrophy, optic atrophy/arterial attenuation

#### TABLE 1. Continued

No.	Individual ID	Haplotype in <i>trans</i>	Age, y First Visit	Sex	Phenotype – Clinical Features
51	Fam11-1	G-A-G	65	F	PD - atrophic patches in foveal region with yellow flecks extending to arcades
52	Fam11-2	G-A-G	67	М	PD - RPE atrophy confined to macular area, surrounded by flecks
53	Fam14-1	G-A-G	69	М	CACD - central RPE and choroidal atrophy
54	Fam14-17	G-A-G	69	М	RP - central RPE and choroidal atrophy and peripheral bone spicules
55	Fam13-60	C-A-A	70	М	PD - butterfly lesion in macula and scattered lesions along the arcades
56	Fam6-5809	G-A-G	70	М	CACD - central RPE and choroidal atrophy
57	Fam13-93	C-A-A	71	F	PD - over 17 y changes from macular lesion to central atrophy
58	Fam13-15	G-A-G	75	F	CRD - RPE atrophy macula and peripapillary region along with bone spicules
59	Fam3-5097	G-A-G	76	F	RP - peripheral bone spicules
60	Fam1-178	G-A-G	84	М	RP - extensive photoreceptor loss, RPE and choroidal atrophy along with bone spicules
61	Fam15-14	C-G-A	85	М	CRD - atrophy and drusen, lots of hyperplasia, cobblestone, geographic atrophy
62	Fam13-33	C-A-A	90	F	PD – peripheral lipofuscin deposits, peripapillary atrophy and scarring, as well as geographic atrophy in macula

The age, sex, and fundus appearance of the affected individuals harboring the splice site mutation, and haplotypes in *trans*,  $Glu_{304}$ -Lys<sub>310</sub>-Gly<sub>338</sub> ( $G_{910}$ -A<sub>929</sub>-G<sub>1014</sub>),  $Gln_{304}$ -Agg<sub>310</sub>-Asp<sub>338</sub> ( $G_{910}$ -G<sub>929</sub>-A<sub>1014</sub>), and  $Gln_{304}$ -Lys<sub>310</sub>-Asp<sub>338</sub> ( $C_{910}$ -A<sub>929</sub>-A<sub>1014</sub>) in exon 3 of *PRPH2* are shown. Classification of the individuals into three phenotypes, PD, CACD, and RP, was based on fundus appearance and other visual function data including ERG and visual fields when available. Min, minimum.



**FIGURE 3.** Visual function in patients with PD (*solid green circles; solid regression line*) and widespread dystrophy (*open red circles*). (A) Visual acuity (decimal) as a function of age. (B) Final dark-adapted threshold as a function of age. *Dashed line* is upper limit of normal. (C) Cone b-wave amplitude to ISCEV standard 30-Hz flicker. *Lines* connect repeated measurements in a patient. *Solid regression* is linear fit to amplitudes for PD. *Dashed line* is lower limit of normal (P < 0.05). (D) Rod b-wave amplitude to ISCEV standard flash. *Lines* connect repeated measurements in a patient. *Solid regression* is linear fit to amplitudes for PD. *Dashed line* is lower limit of normal (P < 0.05).

 TABLE 2.
 Mean and Standard Deviations of ERG Values for Affected (Aff) Individuals Harboring the c.828+3A>T Splice Site Mutation and Normal (Nor) Controls Are Shown

	Aff	Nor	Aff	Nor	Aff	Nor	Aff	Nor
Parameters	Mean	Mean	SD	SD	Min	Min	Max	Max
Age at exam	48		14		20		84	
Rod B amp	50	114	39	24	0	69	187	145
Rod B IT	88	85	9	4	72	79	108	93
Max com amp	193	321	107	37	0	257	420	363
Max com IT	42	39	8	2	31	36	57	42
Cone 30 Hz amp	34	55	21	12	0	34	73	69
Cone 30 Hz IT	34	30	6	1	26	29	54	32
SF cone amp	76	87	44	13	0	65	159	107
SF cone IT	28	29	5	0	14	28	35	29
Dark adaptometry	3		1		1		7	

Amplitudes (amp) are in microvolts, and implicit times (IT) are in milliseconds.

#### **Visual Fields**

Humphrey visual fields were available in 10 individuals. Individuals with PD had better visual fields than those with CRD, CACD, or RP (Figs. 1, 2). Individuals with vitelliform-like PD had the most well-preserved visual fields (Fig. 1A). Butterfly-type PD (Fig. 1B) and flavimaculatus-like PD (Fig. 1C) showed preserved central and mild to moderate peripheral field loss. The CRD and CACD patients had central and paracentral scotoma (Figs. 2A, 2B) and RP patients had primarily peripheral visual field loss progressing to loss of central visual fields as expected (Fig. 2C).

#### **Dark Adaptometry**

Final dark-adapted thresholds were available from 24 patients (Fig. 3B). The majority of patients with PD (green symbols) had thresholds near the upper limit of normal. Thresholds were elevated approximately 2 to 4 log units above normal in individuals with the RP, CRD, and CACD phenotypes (red symbols).

#### Electrophysiology

Electroretinographic data were available for 25 individuals, with many patients tested on multiple occasions. Summary ERG parameters are shown in Table 2 and Figure 3. In general, cone responses were affected less than rod responses. As shown in Figure 3C, patients with PD showed a gradual loss with age of cone amplitude to 30 Hz flicker. The solid green curve is the regression line for cone amplitudes as a function of age on the first visit only. Lines connecting green symbols show repetitive measures on the same individual, with mild decline evident between visits, especially after 40 years of age. In contrast, red symbols show that patients with CRD, CACD, and RP typically had smaller cone amplitudes than patients with

TABLE 3. Frequency of Haplotypes in trans in the Two Groups

Haplotypes	Group I, PD (%)	Group II; CRD, CACD, & RP (%)	Total (%)
C-A-A	13 (29.6)	0 (0)	13 (21)
G-A-G	31 (70.4)	17 (94.4)	48 (77.4)
C-G-A	0 (0)	1 (5.6)	1 (1.6)
Total	44 (71)	18 (29)	62 (100)

Group I with PD and milder phenotype (44 individuals) and Group II with CRD, RP, and CACD with more severe phenotype (18 individuals). Skew in haplotype with Group I having more C-A-A (29.6% vs. 0%) and Group II having more G-A-G (94.4% vs. 70.4%).

PD. Cone b-wave implicit times showed greater delays in patients with more widespread disease than in patients with PD, where cone implicit times were only slightly delayed.

Figure 3D shows rod amplitudes as a function of age. All individuals with rod amplitudes significantly below normal (P < 0.05), with the exception of one individual with PD, had RP, CRD, or CACD (red symbols). Individuals of various subgroups of PD did not show any characteristic pattern of photoreceptor loss. Two individuals who had normal and near-normal amplitudes had vitelliform-like PD (Fig. 1A, #5790) and butterfly-like PD (Fig. 1B, #214). These findings suggested that, although the distribution of flecks results in a varied fundus appearance, the photoreceptor loss does not correspond to fundus appearance.

Long-term follow-up of some individuals over several years show differences in severity of progression. Patients with PD showed milder progressive vision loss (Fig. 4) than patients with CACD/RP-like phenotype (Fig. 5).

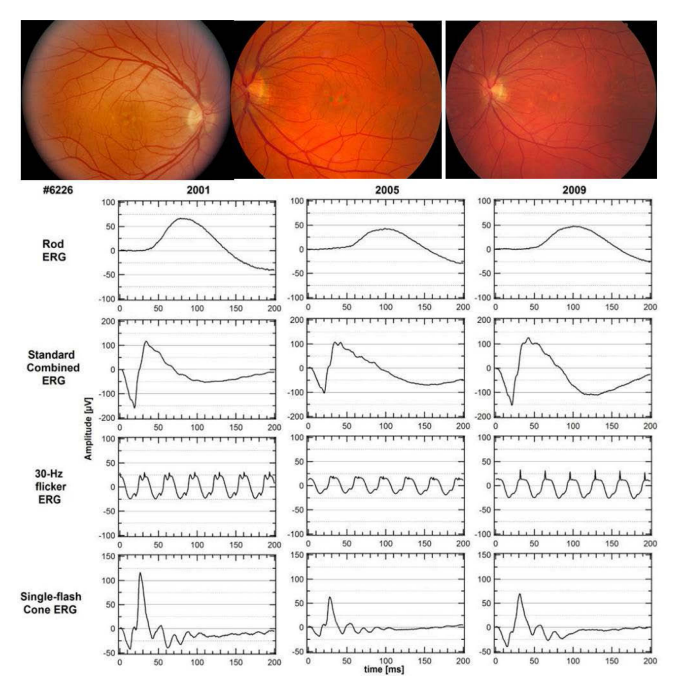
#### **Modifier Gene Screening**

Peripherin/PRPH2. The distribution of PRPH2 (NM\_000322.4) haplotypes in trans for PD, CRD, CACD, and RP phenotypes are shown in Tables 3 and 4. For individuals over 43 years, in Group I, representing milder disease with different types of PDs, Gln<sub>304</sub>-Lys<sub>310</sub>-Asp<sub>338</sub> (C<sub>910</sub>-A<sub>929</sub>-A<sub>1014</sub>) was identified in eight individuals (29.6% vs. 0%) and 19 had Glu<sub>304</sub>-Lys<sub>310</sub>-Gly<sub>338</sub> (G<sub>910</sub>-A<sub>929</sub>-G<sub>1013</sub>). In Group II, with CRD, RP, and CACD, 17 individuals had Glu<sub>304</sub>-Lys<sub>310</sub>-Gly<sub>338</sub> (G<sub>910</sub>-A<sub>929</sub>-G<sub>1013</sub>; 94.4% vs. 70.4%), and one had Gln<sub>304</sub>-Arg<sub>310</sub>-Asp<sub>338</sub>  $(C_{910}-G_{929}-A_{1013})$ .  $Gln_{304}-Lys_{310}-Asp_{338}$   $(C_{910}-A_{929}-A_{1013})$  was absent (Table 4). A significant difference in the haplotype distribution between the two phenotypic groups among individuals older than 43 years was seen using Fisher's exact probability test (P = 0.0092). A GEE analysis that accounted for differences in age and familial correlation of genotypes confirmed that the increased frequency of the Glu<sub>304</sub>-Lys<sub>310</sub>-

**TABLE 4.** Individuals Older than Age 43 Were Used for StatisticalAnalysis and Continue to Exhibit the Skew in Haplotype

Haplotypes	Group I, PD (%)	Group II; CRD, CACD, & RP (%)	Total (%)
C-A-A	8 (29.6)	0 (0)	8 (17.8)
G-A-G	19 (70.4)	17 (94.4)	36 (80.0)
C-G-A	0 (0)	1 (5.6)	1 (2.2)
Total	27 (60)	18 (40)	45 (100)

Fisher, P = 0.009; GEE analysis, 0.03.



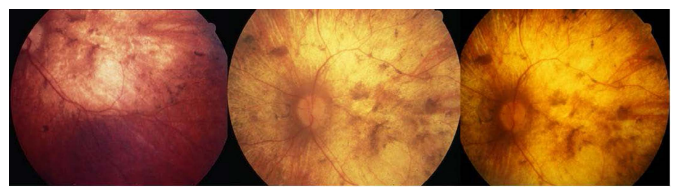
**FIGURE 4.** Pattern dystrophy in a female aged 36 years (Fam8-6226). Milder vision loss and slow progression over a 9-year period, with relatively preserved rod and cone function at age 45. Lower limits of normal (P < 0.05) are 85 µV for rod ERG, 200 µV for standard combined ERG, 50 µV for 30-Hz flicker ERG, and 75 µV for single flash cone ERG.

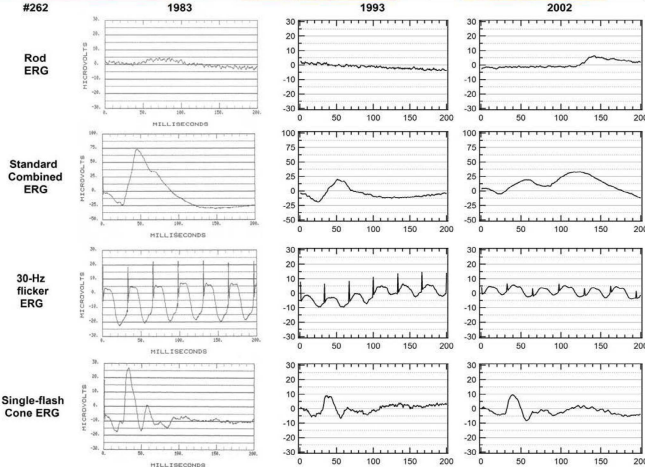
Gly<sub>338</sub> haplotype among individuals with CRD, RP, and CACD was significant with a relevant *P* value of <2e-16. However, because the asymptotic *P* value may not be robust to the "zero" cell counts in two of the haplotype-phenotype categories (see Tables 3, 4), we reran the GEE model taking the conservative approach of collapsing the Gln<sub>304</sub>-Lys<sub>310</sub>-Asp<sub>338</sub> (C-A-A) and Gln<sub>304</sub>-Arg<sub>310</sub>-Asp<sub>338</sub> (C-G-A) haplotypes into a single category (*P* = 0.03, *N* = 45 due to removal of individuals with age <43). The estimated odds ratio for this comparison is 7.16 (95% CI = [2.8-18.4]), suggesting that the odds of having the CACD/RP-like phenotype (versus the PD phenotype) are 7.16 times greater for affected individuals with

a Glu<sub>304</sub>-Lys<sub>310</sub>-Gly<sub>338</sub> haplotype (G-A-G) haplotype in *trans* than for affected individuals with one of the other two haplotypes in *trans* (Gln<sub>304</sub>-Lys<sub>310</sub>-Asp<sub>338</sub> [C-A-A] and Gln<sub>304</sub>-Arg<sub>310</sub>-Asp<sub>338</sub> [C-G-A]).

**ROM1** Gene. No significant sequence variants were identified by bidirectional sequencing of *ROM1* (NM\_000327.3), in 10 affected individuals representing different phenotypes.

*GUCA1A* Gene. A *GUCA1A* (NM\_000409.3), c.149C>T (p.Pro50Leu), sequence variant was identified in 4 individuals (3 affected—1 with macular atrophy and 2 with PD; 1 individual had normal retina) by bidirectional sequencing of *GUCA1A* in 62 individuals.





**FIGURE 5.** Retinitis pigmentosa in a male aged 48 years (Fam1-365). Rapid progression over a 20-year period from relatively preserved photoreceptors and RPE to a complete loss of rod and greatly reduced cone function. Lower limits of normal (P < 0.05) are 85 µV for rod ERG, 200 µV for standard combined ERG, 50 µV for 30-Hz flicker ERG, and 75 µV for single flash cone ERG.

*NXNL1* Gene. Several variants including c.46G>A (p.Asp16Asn), c.83G>C (p.Arg28Pro), c.172\_174delTTC, (p.del-Phe58), c.190G>A (p.Glu64Lys), and c.461A>T (p.Glu154Val), were identified by bidirectional screening of *NXNL1* (NM\_138454.1) in 62 affected individuals and 103 CEPH normal controls. However, none showed any significant association.

### DISCUSSION

This study demonstrated that the c.828+3A>T splice site mutation in the *PRPH2* gene is the primary cause of retinal

disease in all 62 affected individuals in these 19 families with complete penetrance but highly variable expressivity. It results in clinically diverse, autosomal dominant retinal dystrophies, including PDs, CRD, CACD, and RP (Figs. 1, 2), in 19 families derived from a common ancestor.<sup>2</sup> The phenotypes fall into two broad clinical categories: Individuals in Group I (N = 44; AVMD, BMD, and AOFMD types of PD) have milder disease than individuals in Group II (CRD, CACD, and RP). There is limited progression from Groups I to II. In our cohort, almost all individuals had reduced photoreceptor responses relative to age-matched controls, regardless of the phenotype (Fig. 3).

We previously had established a founder effect for the *PRPH2* c.828+3A>T mutation by haplotype analysis which

excludes sequence variants in cis within 5 megabases of the PRPH2 locus as modifiers.<sup>2</sup> In this study, we analyzed the haplotypes in trans and showed that Gln<sub>304</sub>-Lys<sub>310</sub>-Asp<sub>338</sub> (C<sub>910</sub>-A<sub>929</sub>-A<sub>1014</sub>) is significantly associated with Group I pattern dystrophies and Glu<sub>304</sub>-Lys<sub>310</sub>-Gly<sub>338</sub> (G<sub>910</sub>-A<sub>929</sub>-G<sub>1013</sub>) with Group II (P value = 0.03 taking family structure, progression with age and restricting our analysis to only individuals (N = 45) older than 43 years of age. This suggested the possible modifier role of the polymorphic protein haplotypes in trans, with an estimated 7.16-fold increase in the odds of the more severe phenotype for affected individuals harboring a Glu304-Lys310-Gly338 (G-A-G) haplotype in trans compared to other affected individuals, and a possible protective role for having a second Gln<sub>304</sub>-Lys<sub>310</sub>-Asp<sub>338</sub> (C-A-A) haplotype in *trans*. Given that the two alleles are similar in their temporal and spatial expression and interact directly, the protein haplotype in trans is likely to contribute to the phenotypic variation seen in the PRPH2 splice site mutation families.

Several sequence variants were found in the other potential modifier genes that were screened; however, none of them had a significant association with any one phenotype. Progressive photoreceptor loss with age explains continued vision loss in some individuals. Another possible modifier may be variations in expression of normal versus aberrant transcripts due to leaky aberrant splicing.<sup>2</sup> Functional studies with more accurate quantification of the truncated protein and its interaction with different RDS protein mini-haplotypes may help determine the effect of variable splicing and protein mini-haplotypes in modifying the retinal phenotype. Future studies with larger number of patients and whole exome sequencing of additional retinal disease-causing genes and/or whole exome and genome sequencing may identify additional modifiers.

This study demonstrated the variability of autosomal dominant retinal dystrophies resulting from *PRPH2* mutations and the possibility of common polymorphic protein variants functioning as modifiers of disease-causing mutations in Mendelian disorders. Determining the protein haplotype in *trans* in these families may help predict the severity of the retinal phenotype and may aid in counseling of the families. The potential for splice site modification also is of therapeutic interest in preserving vision in the affected individuals and needs further evaluation.

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