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

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PRPH2 mutation update: In silico assessment of 245 reported and 7 novel variants in patients with retinal disease

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Abstract

Mutations in *PRPH2*, encoding peripherin-2, are associated with the development of a wide variety of inherited retinal diseases (IRDs). To determine the causality of the many *PRPH2* variants that have been discovered over the last decades, we surveyed all published *PRPH2* variants up to July 2020, describing 720 index patients that in total carried 245 unique variants. In addition, we identified seven novel *PRPH2* variants in eight additional index patients. The pathogenicity of all variants was determined using the ACMG guidelines. With this, 107 variants were classified as pathogenic, 92 as likely pathogenic, one as benign, and two as likely benign. The remaining 50 variants were classified as variants of uncertain significance. Interestingly, of the total 252 *PRPH2* variants, more than half ($n = 137$) were missense variants. All variants were uploaded into the Leiden Open source Variation and ClinVar databases. Our study underscores the need for experimental assays for variants of unknown significance to improve pathogenicity classification, which would allow us to better understand genotype-phenotype correlations, and in the

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long-term, hopefully also support the development of therapeutic strategies for patients with *PRPH2*-associated IRD.

KEYWORDS

in silico assessment, inherited retinal disease, LOVD, molecular genetics, *PRPH2*

1 | INTRODUCTION

PRPH2, also known as retinal degeneration slow (RDS), was first described in 1991 (Kajiwara et al., 1991). The gene encodes peripherin-2, a 346 amino-acid long glycoprotein that spans the membrane four times, and is located primarily in the rim regions of rod and cone outer segment (OS) discs and lamellae. Besides the four transmembrane domains, the protein contains a cytoplasmic (C) loop and two intra-discal (D1 and D2) loops (Travis et al., 1989, 1991, 1992). Its exact molecular function inside photoreceptor cells is not yet fully understood, but it is hypothesized that the protein plays an essential role in the initiation of OS disc formation, as well as in disc stabilization, maintenance, and disc size alignment, mainly by forming oligomers with other *PRPH2* molecules and/or Retinal Outer Segment Membrane Protein 1 (ROM1) (Chakraborty et al., 2020; Zulliger et al., 2018). For instance, *Prph2*^{-/-} mice failed to initiate OS disc formation, whereas *Prph2* p.Cys150-Ser^{+/-} mice did not support proper OS formation, interacted abnormally with Rom1, and showed reduced *Prph2* protein levels (Zulliger et al., 2018). In addition, it was shown that in *Prph2* p.Cys213Tyr^{-/-} mice, mutant *Prph2* could not oligomerize with ROM1 and was mislocalized, being retained in the inner segments (Chakraborty et al., 2020). Based on these studies, *PRPH2* indeed seems to be critical for proper OS formation as well as for its function.

To date, over 200 different *PRPH2* variants have been described to be associated with the development of a wide variety of inherited retinal diseases (IRDs), such as retinitis pigmentosa (RP), cone-rod dystrophy (CRD), and macular dystrophies. The group of *PRPH2*-associated macular dystrophies encompasses a wide variety of phenotypes, including pseudo-Stargardt pattern dystrophy, butterfly-shaped pigment dystrophy (BPD), adult-onset foveomacular vitelliform dystrophy (AOFVD), and central areolar choroidal dystrophy (CACD) (Boon et al., 2008; 2009; Boon, van Schooneveld, et al., 2007; Kersten et al., 2018). These macular dystrophy phenotypes, especially CACD, may be confused with geographic atrophy in age-related macular degeneration (AMD), and *PRPH2* mutations have been described in patients initially diagnosed with AMD (Boon et al., 2009; Kersten et al., 2018; Smailhodzic et al., 2011). *PRPH2* mutations are most frequently inherited in an autosomal dominant fashion, although autosomal recessive and sporadic cases have also been reported, as well as autosomal dominant cases with reduced penetrance (Alapati et al., 2014; Birtel et al., 2018; Boon et al., 2008, 2009; Coco et al., 2010; Dryja et al., 1997; Khan et al., 2016; Manes et al., 2015). Interestingly, heterozygous mutations in both *PRPH2* and *ROM1* can cause digenic RP (Kajiwara et al., 1994).

In this study, we performed a systematic analysis of all 245 *PRPH2* variants in IRD patients that were published to date. For this purpose, we collected all *PRPH2* variants published up to July 2020 that were associated with the development of IRD. In addition, we added *PRPH2* variants that were either identified via our routine diagnostics pipeline, or via a novel sequencing approach using molecular inversion probes (MIPs) (Hiatt et al., 2013; Neveling et al., 2017; Weisschuh et al., 2018), seven of which have not been described before. All variants were classified for their pathogenicity using the American College of Medical Genetics and Genomics (ACMG) guidelines, after which they were uploaded into the Leiden Open source Variation Database (LOVD) and ClinVar database for *PRPH2*. In addition, we attempted to establish genotype-phenotype correlations. Several variants are reported to be associated with only one specific IRD phenotype, and these diseases can best be described as allelic disorders. However, for most variants in this study, there appears to be a high phenotypic variability between different families, as well as between members of the same family.

By performing this study, we aim to shed light on how to experimentally assess the true causality of *PRPH2* variants in the future, as well as to explain the observed phenotypic variability within IRD patients. This will facilitate a better interpretation of the pathogenicity of variants that are identified in subjects with IRD, and in the long term, hopefully also support the development of therapeutic strategies for patients with *PRPH2*-associated IRD.

2 | MATERIAL AND METHODS

2.1 | Literature search

We collected all publications up to July 2020 that report *PRPH2* variants in patients with inherited retinal disease. The following PubMed search terms were used: "(retinal+degeneration+slow+OR+PRPH2+OR+peripherin)+AND+(central+areolar+choroidal+dystrophy+OR+caccd+OR+vision+disorders+OR+retinal+dystrophy)+AND+(mutation+OR+variant+OR+mutations+OR+variants)". In addition, the HGMD professional database was used to search for variants or articles that were possibly missed with our PubMed queries. Variant detection, variant combinations, patient age, patient gender and age at onset, disease phenotype, segregation analysis, and allele frequencies (gnomAD) were collected. Obvious duplicates, in some cases following contact with the corresponding authors of the respective papers, were removed from the data set. An important point to take into consideration when retracting genetic variants from literature search tools is that it is always possible to miss certain variants. For instance, a *PRPH2* variant might be present in

Supporting Information data of papers that could be potentially missed by both PubMed and HGMD searches.

2.2 | Subjects

This study was approved by the institutional review boards of the Radboud University Medical Center (Radboudumc) and was conducted in adherence to the tenets of the Declaration of Helsinki. All Sanger sequencing and/or whole (exome) sequencing data from the Radboudumc genome diagnostic laboratory were analyzed to determine the causative genetic defects in patients with visual impairment. *PRPH2* variants were also identified by using a targeted sequencing approach based on molecular inversion probes (MIPs) (Hiatt et al., 2013; Neveling et al., 2017; Weisschuh et al., 2018). Single-molecule MIPs were synthesized to capture and sequence overlapping 110-nt segments of the three *PRPH2* protein-coding exons and flanking splice sites, similarly as described previously for the *ABCA4* gene (Khan et al., 2020). *PRPH2* variants identified with this MIPs approach were validated with Sanger sequencing.

2.3 | Variant analysis

The complementary DNA (cDNA) was numbered as follows: the A of the ATG translation initiation codon of the *PRPH2* reference sequence (NM_000322.5) was reported as +1 while the initiation codon was reported as codon 1. Allele frequencies of *PRPH2* variants in control populations were extracted from the genome aggregation database (gnomAD: v2.1.1 and v3). This database contains both whole exome and whole genome sequencing data obtained from 213,158 healthy individuals from all over the world (<https://gnomad.broadinstitute.org/>).

Next, we performed statistical analysis to compare allele frequencies in the index patient group to the normal population (gnomAD). This enabled us to assess whether a specific variant is enriched in patient versus control groups. For this purpose, the Fisher's Exact test was used. To only select truly statistically significant findings, a correction by the false discovery rate (FDR) of Benjamini-Hochberg, classical one stage method with an error margin of 5%, was carried out (Benjamini et al., 2001).

2.4 | In silico predictions

For *PRPH2* missense variants, Polyphen-2 (benign = 0.000–0.150; possibly damaging = 0.150–0.850; probably damaging = 0.850–1.000), Combined Annotation Dependent Depletion (CADD, tolerated = <15; damaging = >15), and SIFT scores (tolerated = 0.050–1.000; damaging = 0.000–0.050) were obtained from <http://genetics.bwh.harvard.edu/pph2/>, <http://cadd.gs.washington.edu/home> (Kircher et al., 2014) and <https://sift.bii.a-star.edu.sg/>, respectively. For splice site variants, software to gather splicing scores available via Alamut Visual version 2.13 (Interactive Biosoftware) was used.

2.5 | Variant pathogenicity classification

Pathogenicity of all reported and newly identified *PRPH2* variants was predicted using the ACMG guidelines (Richards et al., 2015). These guidelines enabled us to classify each variant into one of the following categories: pathogenic, likely pathogenic, benign, likely benign, or of uncertain significance. To do so, we scored the variants based on the evidence of pathogenicity in different ACMG categories:

1. Pathogenic, very strong (PVS), for example, this variant is protein-truncating.
2. Pathogenic, strong (PS), for example, this variant leads to the same amino acid being substituted compared to a previously described pathogenic variant.
3. Pathogenic, moderate (PM), for example, this variant is located in a mutational hot spot and/or well-established functional domain.
4. Pathogenic, supporting (PP), for example, multiple lines of computational evidence support the variant to be pathogenic.

The scores of each category are combined to come to the final classification in one of the five pathogenicity categories that are explained in more detail in Table S1.

2.6 | LOVD and ClinVar submission

The 245 reported, as well as the seven novel *PRPH2* variants were uploaded to the LOVD and ClinVar database, when available, together with patient data including the description of the phenotype, patient age, patient age of onset, and segregation information. Pathogenicity scores of all variants were based on the pathogenicity assessment as described in the "In silico predictions" section.

3 | PRPH2 VARIANTS

3.1 | PRPH2 mutation spectrum

We collected 245 *PRPH2* variants identified in 720 index patients that were described in 165 articles up to July 2020, as well as their phenotypic information (Abouelhoda et al., 2016; Abu-Safieh et al., 2013; Ahmad et al., 2010; Alapati et al., 2014; Anand et al., 2009; Anasagasti et al., 2013; Apfelstedt-Sylla et al., 1995; Arai et al., 2015; Avela et al., 2019, 2018; Ba-Abbad et al., 2019, 2014; Barbazetto et al., 2007; Bareil et al., 2000, 1997; Bayés et al., 1996; Birtel et al., 2018; Boon et al., 2009; Boon, Klevering, et al., 2007; Boon, van Schooneveld, et al., 2007; Boulanger-Scemama et al., 2015; Budu et al., 2001; Carss et al., 2017; Cheng et al., 2019; Coco-Martin et al., 2020; Coco et al., 2010; Consugar et al., 2015; Coussa et al., 2015; Daftarian et al., 2019; de Breuk et al., 2020; Deciphering Developmental Disorders Study, 2017; de Sousa Dias et al., 2015; Donoso et al., 2003; Downes et al., 1999; Downs et al., 2007; Dryja et al., 1997; Duncan et al., 2011; Duncker et al., 2015; Ekström, Andréasson, et al., 1998;

Ekström, Ponjavic, et al., 1998; Ekström, Ponjavic, Andréasson, et al., 1998; Essilfie et al., 2018; Fakin et al., 2012; Farrar et al., 1992, 1991; Feist et al., 1994; Felbor et al., 1997; Fernandez-San Jose et al., 2015; Fishman et al., 1994, 1997; Foote et al., 2019; Francis et al., 2005; Gamundi et al., 2007; Gao et al., 2019; Glöckle et al., 2014; Gocho et al., 2016; Gorin et al., 1995; Grover et al., 2002; Grüning et al., 1994; Hanany & Sharon, 2019; Hosono et al., 2018; Hoyng et al., 1996; Huang et al., 2013; Jacobson et al., 1996, 2016; Jespersgaard et al., 2019; Jin et al., 2008; Jones et al., 2017; Kajiwara et al., 1994, 1991, 1993; Kalyanasundaram et al., 2009; Katagiri et al., 2018; Keen et al., 1994; Keilhauer, Meigen, Stöhr, et al., 2006; Keilhauer, Meigen, & Weber, 2006; Kemp et al., 1994; Kersten et al., 2018; Khan, 2019; Khan et al., 2016; Khoubian et al., 2005; Kikawa et al., 1994; C. Kim et al., 2012; R. Y. Kim et al., 1995; Kitiratschky et al., 2011; Klevering et al., 2002; Kohl et al., 1997, 2012; Lam et al., 1995; C. S. Lee & Leys, 2020; K. Lee et al., 2015; Leroy et al., 2007; Lim et al., 2009; Maertz et al., 2015; Manes et al., 2015; Martin-Merida et al., 2018; Meins et al., 1993; Meunier et al., 2011; Michaelides et al., 2005; Miyata et al., 2018; Moshfeghi et al., 2006; Nakazawa et al., 1994, 1995, 1996; Nanda et al., 2019; Neveling et al., 2012; Nichols, Drack, et al., 1993; Nichols, Sheffield, et al., 1993; Oishi et al., 2014; Pajic et al., 2006; Palma et al., 2019; Passerini et al., 2007; Patel et al., 2016; Payne et al., 1998; Poloschek et al., 2010; Ramkumar et al., 2017; Reeves et al., 2020; Reig et al., 1995; Renner et al., 2004, 2009; Richards & Creel, 1995; Saga et al., 1993; Sallevelt et al., 2017; Schatz et al., 2003; Schorderet et al., 2014; Sears et al., 2001; Shankar et al., 2015, 2016; Simonelli et al., 2007; Smailhodzic et al., 2011; Sohocki et al., 2001; Souied et al., 1998; Stone et al., 2017; Strafella et al., 2019; Strom et al., 2012; Sullivan et al., 2006, 2013; Sun et al., 2015; Testa et al., 2005; Trujillo et al., 1998, 2001; Trujillo Tiebas et al., 2002; Vaclavik et al., 2012; Van Cauwenbergh et al., 2017; van Lith-Verhoeven et al., 2003; X. Wang et al., 2013; J. Wang et al., 2014; H. Wang et al., 2015; Wawrocka et al., 2018; Weleber et al., 1993; Wells et al., 1993; Wolock et al., 2019; Wroblewski et al., 1994a; 1994b; Xiang et al., 2012; Xu et al., 2014; Xue et al., 2014; Yanagihashi et al., 2003; H. Yang et al., 2000; Z. Yang et al., 2003; Z. Yang et al., 2004; Yeoh et al., 2010; Yoon et al., 2015; Zaneveld et al., 2015; Zhang et al., 2002; Zhao et al., 2015; Zhou et al., 2018; Zhuk & Edwards, 2006). Additionally, 139 index cases from either the Radboudumc genome diagnostic laboratory or that were studied via MIPs analysis, carried *PRPH2* variants that were published before (Table 1). Finally, we also identified seven novel variants in eight additional index patients that, to our knowledge, were never identified (Table 1; variants depicted in bold lettering). Of the collective 252 *PRPH2* variants, 137 were missense, 85 were protein-truncating, 10 were splice sites, 15 were in-frame amino acid changes, three were synonymous, and two were located in the 5'- or 3'-untranslated regions (UTRs) (Figure S1A; Table 2). Of the total 720 previously reported index patients, 686 patients carried heterozygous variants, 8 patients carried compound heterozygous variants, 14 patients carried homozygous variants, and 11 patients had variants in two different genes. There was one patient that carried variants in three different genes (*PRPH2*, *ROM1*, *ABCA4*). The authors speculated trigenic

inheritance; however, it still needs to be elucidated to determine whether this case is truly trigenic. The first homozygous *PRPH2* variants were identified by Wang and colleagues. A cohort of 179 index cases was genetically screened leading to the identification of two unrelated patients carrying the homozygous p.Leu185Pro and p.Cys213Trp *PRPH2* variants, respectively (X. Wang et al., 2013). These patients were both suffering from Leber congenital amaurosis (LCA), suggesting that homozygous *PRPH2* variants are associated with more severe phenotypes compared to heterozygous *PRPH2* mutation carriers. Of the 139 index cases identified by the Radboudumc genome diagnostic laboratory or our MIPs analysis, two patients also carried homozygous variants, while the remaining 137 index cases were all heterozygous carriers. The two index cases that were found to be homozygous both carried the *PRPH2*-p.Arg142Trp variant, and were diagnosed with CACD. To our knowledge, this is the first time this particular variant was present in a homozygous state. For one of the patients, we were able to genetically screen both parents. The father was also affected by CACD. Interestingly, the mother did not show apparent signs of retinal disease upon ophthalmological examination. Both parents were proven to carry the variant in a heterozygous state. When studying the phenotype of both patients in more detail, we could observe that it resembles the phenotype we observe in heterozygous p.Arg142Trp carriers, pointing toward a semi-dominant inheritance with a dominant-negative effect on the wildtype allele. Finally, the eight index patients carrying the seven novel *PRPH2* variants, were all heterozygous for the respective variants. All variants, together with a description of the phenotype and, when available, segregation analysis, were uploaded to the LOVD and ClinVar database (<https://databases.lovd.nl/shared/genes/PRPH2>).

3.2 | Recurrent *PRPH2* variants

The most recurrent *PRPH2* variant among IRD patients is p.Arg142Trp. This variant was reported in 95 out of 867 index patients, of which 93 were heterozygous carriers and two were homozygous (Table S2). The variant was exclusively reported in individuals with Caucasian ancestry, and in the Netherlands, this variant is mainly associated with the development of CACD. Analysis of single nucleotide polymorphisms (SNPs) in close proximity of the p.Arg142Trp variant in three Dutch families revealed the presence of a shared chromosomal segment of approximately 519 kb, strongly suggesting that this particular variant represents a founder mutation in the Netherlands (Boon et al., 2009). As mentioned in the introduction section, CACD may be confused with AMD, and besides retinal imaging modalities (optical coherence tomography and fundus autofluorescence), screening for the p.Arg142Trp may help to better discriminate between these two phenotypes (Kersten et al., 2018; Smailhodzic et al., 2011). One interesting observation in large Dutch CACD families harboring this variant is that there seems to be reduced penetrance (Boon et al., 2009). Within these families, some individuals were significantly less severely affected compared to age-matched family members also carrying the variant. The molecular mechanisms behind this phenomenon still need to be elucidated. Possible explanations could be:

TABLE 1 Previously unreported *PRPH2* variants identified in index cases by MIPs or the Radboudumc genome diagnostic laboratory

PID	cDNA change	Protein change	Type of mutation	Identified by	Phenotype
1	c.2T>C	p.?	Fail-to-start	MIPs	Pseudo-STGD
2	c.63G>A	p.Trp21*	Nonsense	Exome seq.	MD/PD
3	c.63G>A	p.Trp21*	Nonsense	Sanger seq.	MD/PD
4	c.63G>A	p.Trp21*	Nonsense	Sanger seq.	MD/PD
5	c.94A>G	p.Ile32Val	Missense	MIPs	AMD
6	c.112G>T	p.Gly38*	Nonsense	MIPs	Pseudo-STGD
7	c.122T>C	p.Leu41Pro	Missense	Sanger seq.	MD/PD
8	c.133C>T	p.Leu45Phe	Missense	MIPs	AMD
9	c.136C>T	p.Arg46*	Nonsense	MIPs	Pseudo-STGD
10	c.209dup	p.Ser71Ilefs*106	Frameshift	MIPs	Pseudo-STGD
11	c.253G>A	p.Ala85Thr	Missense	Sanger seq.	MD/PD
12	c.271T>A	p.Tyr91Asn	Missense	MIPs	Pseudo-STGD
13	c.281G>A	p.Trp94*	Nonsense	MIPs	Pseudo-STGD
14	c.303C>G	p.Tyr101*	Nonsense	Exome seq.	MD/PD
15	c.367C>T	p.Arg123Trp	Missense	Exome seq.	RP
16	c.377T>C	p.Leu126Pro	Missense	Exome seq.	RP
17	c.415_417del	p.Lys139del	Deletion	MIPs	Pseudo-STGD
18	c.415_417del	p.Lys139del	Deletion	MIPs	Pseudo-STGD
19	c.423C>A	p.Tyr141*	Nonsense	MIPs	Pseudo-STGD
20	c.424C>T	p.Arg142Trp	Missense	MIPs	Pseudo-STGD
21	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
22	c.424C>T	p.Arg142Trp	Missense	Exome seq.	RD
23	c.424C>T	p.Arg142Trp	Missense	Exome seq.	RCD
24	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
25	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
26	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
27	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
28	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
29	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
30	c.424C>T	p.Arg142Trp	Missense	Exome seq.	CACD
31	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD/CD
32	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
33	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD/PD
34	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
35	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
36	c.424C>T	p.Arg142Trp	Missense	Exome seq.	CACD
37	c.424C>T	p.Arg142Trp	Missense	Exome seq.	MD
38	c.424C>T	p.Arg142Trp	Missense	Exome seq.	RD
39	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD

(Continues)

TABLE 1 (Continued)

PID	cDNA change	Protein change	Type of mutation	Identified by	Phenotype
40	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
41	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	MD/PD
42	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
43	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
44	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
45	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
46	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
47	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	MD/PD
48	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	MD/PD
49	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
50	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
51	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
52	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	MD/PD
53	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
54	c.424C>T	p.Arg142Trp	Missense	Sanger seq.	CACD
55	c.433_434del	p.Asp145Hisfs*31	Frameshift	Sanger seq.	CACD
56	c.441del	p.Gly148Alafs*5	Frameshift	MIPs	Pseudo-STGD
57	c.441del	p.Gly148Alafs*5	Frameshift	MIPs	Pseudo-STGD
58	c.441del	p.Gly148Alafs*5	Frameshift	Sanger seq.	MD/PD
59	c.441del	p.Gly148Alafs*5	Frameshift	Sanger seq.	MD/PD
60	c.441del	p.Gly148Alafs*5	Frameshift	Exome seq.	MD
61	c.458A>G	p.Lys153Arg	Missense	Asper	RP
62	c.469G>A	p.Asp157Asn	Missense	MIPs	Pseudo-STGD
63	c.499G>A	p.Gly167Ser	Missense	Sanger seq.	MD/PD
64	c.499G>A	p.Gly167Ser	Missense	MIPs	Pseudo-STGD
65	c.505_507del	p.Asn169del	Deletion	Sanger seq.	MD/PD
66	c.505_507del	p.Asn169del	Deletion	Exome seq.	RP
67	c.514C>T	p.Arg172Trp	Missense	MIPs	Pseudo-STGD
68	c.514C>T	p.Arg172Trp	Missense	MIPs	Pseudo-STGD
69	c.514C>T	p.Arg172Trp	Missense	MIPs	Pseudo-STGD
70	c.514C>T	p.Arg172Trp	Missense	Asper	RD
71	c.514C>T	p.Arg172Trp	Missense	Sanger seq.	MD/PD
72	c.514C>T	p.Arg172Trp	Missense	Sanger seq.	MD/PD
73	c.514C>T	p.Arg172Trp	Missense	Exome seq.	CD
74	c.514C>T	p.Arg172Trp	Missense	Exome seq.	MD
75	c.514C>T	p.Arg172Trp	Missense	Exome seq.	MD
76	c.515G>A	p.Arg172Gln	Missense	MIPs	Pseudo-STGD
77	c.515G>A	p.Arg172Gln	Missense	Exome seq.	MD
78	c.515G>A	p.Arg172Gln	Missense	MIPs	Pseudo-STGD

PID	cDNA change	Protein change	Type of mutation	Identified by	Phenotype
79	c.520T>A	p.Trp174Arg	Missense	Sanger seq.	MD/PD
80	c.520T>A	p.Trp174Arg	Missense	Exome seq.	MD
81	c.522G>C	p.Trp174Cys	Missense	MIPs	Pseudo-STGD
82	c.581+1G>A	p.?	Splice site	MIPs	Pseudo-STGD
83	c.581+4dup	p.?	Splice site	Sanger seq.	?
84	c.582-1G>A	p.?	Splice site	MIPs	Pseudo-STGD
85	c.582-1G>A	p.?	Splice site	Sanger seq.	CACD
86	c.582-2A>T	p.?	Splice site	Sanger seq.	CACD
87	c.582_828del	p.Asp194Glufs*2	Frameshift	MIPs	Pseudo-STGD
88	c.583C>T	p.Arg195*	Nonsense	MIPs	Pseudo-STGD
89	c.583C>T	p.Arg195*	Nonsense	MIPs	Pseudo-STGD
90	c.584G>T	p.Arg195Leu	Missense	MIPs	Pseudo-STGD
91	c.584G>A	p.Arg195Gln	Missense	Sanger seq.	CACD
92	c.584G>A	p.Arg195Gln	Missense	MIPs	Pseudo-STGD
93	c.614T>C	p.Leu205Pro	Missense	Sanger seq.	CACD
94	c.623G>A	p.Gly208Asp	Missense	MIPs	Pseudo-STGD
95	c.623G>A	p.Gly208Asp	Missense	Sanger seq.	MD/PD
96	c.623G>A	p.Gly208Asp	Missense	Sanger seq.	MD/PD
97	c.623G>A	p.Gly208Asp	Missense	MIPs	AMD
98	c.628C>T	p.Pro210Ser	Missense	MIPs	Pseudo-STGD
99	c.633C>A	p.Phe211Leu	Missense	Asper	RP
100	c.646C>T	p.Pro216Ser	Missense	Asper	RP
101	c.646C>T	p.Pro216Ser	Missense	Exome seq.	RCD
102	c.646C>T	p.Pro216Ser	Missense	Exome seq.	RP
103	c.646C>T	p.Pro216Ser	Missense	Exome seq.	RP
104	c.646C>T	p.Pro216Ser	Missense	Sanger seq.	RP
105	c.646C>T	p.Pro216Ser	Missense	Exome seq.	RP
106	c.656C>G	p.Pro219Arg	Missense	Sanger seq.	CACD
107	c.657_662del	p.Arg220_Pro221del	Deletion	Sanger seq.	MD/PD
108	c.658C>T	p.Arg220Trp	Missense	Sanger seq.	MD/PD
109	c.658del	p.Arg220fs*34	Frameshift	Exome seq.	RCD/CRD
110	c.658del	p.Arg220fs*34	Frameshift	Sanger seq.	MD/PD
111	c.659G>A	p.Arg220Gln	Missense	Sanger seq.	MD/PD
112	c.665G>A	p.Cys222Try	Missense	MIPs	CRD
113	c.715C>T	p.Gln239*	Missense	MIPs	AMD
114	c.736T>C	p.Trp246Arg	Missense	MIPs	AMD
115	c.746del	p.Gly249Alafs*7	Frameshift	Exome seq.	MD/PD
116	c.749G>A	p.Cys250Tyr	Missense	MIPs	Pseudo-STGD
117	c.754G>C	p.Ala252Pro	Missense	MIPs	AMD

(Continues)

TABLE 1 (Continued)

PID	cDNA change	Protein change	Type of mutation	Identified by	Phenotype
118	c.781C>T	p.Leu261Phe	Missense	MIPs	AMD
119	c.809_810del	p.Leu270Profs*30	Frameshift	MIPs	Pseudo-STGD
120	c.850C>T	p.Arg284Cys	Missense	MIPs	AMD
121	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
122	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
123	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
124	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
125	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
126	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
127	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
128	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
129	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
130	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
131	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
132	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
133	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
134	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
135	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
136	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
137	c.866C>T	p.Ser289Leu	Missense	MIPs	AMD
138	c.866C>T	p.Ser289Leu	Missense	Sanger seq.	MD/PD
139	c.923T>A	p.Leu308Gln	Missense	MIPs	AMD
140	c.923T>A	p.Leu308Gln	Missense	MIPs	AMD
141	c.938C>T	p.Pro313Leu	Missense	MIPs	AMD
142	c.938C>T	p.Pro313Leu	Missense	MIPs	AMD
143	c.938C>T	p.Pro313Leu	Missense	MIPs	AMD
144	c.828+1G>A	p.?	Splice site	Sanger seq.	MD/PD
145	c.829-3_829-1del	p.?	Splice site	MIPs	Pseudo-STGD
146	c.897_898del	p.Ser301Argfs*90	Frameshift	Exome seq.	MD
147	c.946T>G	p.Trp316Gly	Missense	Exome seq.	MD/PD

Note: Novel variants identified via MIPs are depicted in bold.

Abbreviations phenotypes: AMD, age-related macular degeneration; CACD, central areolar choroidal dystrophy; CD, cone dystrophy; CRD, cone-rod dystrophy; MD, macular dystrophy; PD, pattern dystrophy; pseudo-STGD, pseudo-Stargardt disease; RD, retinal dystrophy; RP, retinitis pigmentosa; RCD, rod-cone dystrophy.

(1) reduced expression of the mutant allele; (2) increased expression of the wild-type allele; (3) the influence of modifier alleles located in other genes; or (4) influence of environmental factors (e.g., smoking, nutrition).

Other recurrently reported variants (reported in >10 index patients) are p.Arg46*, p.Tyr141Cys, p.Gly148Alafs*5, p.Arg172Trp, p.Arg172Gln, p.Leu185Pro, p.Arg195Leu, p.Gly208

Asp, p.Pro210Arg, p.Pro216Ser, p.Pro216Leu, p.Gln239*, p.Ser289Leu, and c.828+3A>T. The p.Arg172Trp variant was reported in 60 index patients and was mainly found in Caucasians with British ancestry. There was only one Japanese index patient carrying this variant. Of the total 60 cases, 58 were heterozygous for this variant (Table S2). Payne and colleagues performed

TABLE 2 Distribution of *PRPH2* variants found in IRD patients

Variant type	Number of unique variants	Number of alleles
Missense	137	605
Protein-truncating	85	180
Splice site	10	77
In-frame amino acid insertions/deletions	15	24
Synonymous	3	3
5'- or 3'-UTR	2	2
Total	252	891

Abbreviation: UTR, untranslated region.

haplotype analysis in multiple British families carrying the stand-alone p.Arg172Trp variant and revealed that in Britain, this is a founder mutation (Payne et al., 1998). A German index patient carried an additional pathogenic heterozygous *ROM1* missense variant (p.Arg229His), and another patient carried two additional heterozygous missense variants, one in *ROM1* (p.Arg229His) and one in *ABCA4* (p.Val2050Leu (Table S2) (Poloschek et al., 2010). The authors argued that the *ROM1* and *ABCA4* variants act as a moderator, worsening the pattern dystrophy phenotype compared to individuals that only carry the p.Arg172Trp variant in *PRPH2*. However, when applying the ACMG criteria to both the *ROM1* and *ABCA4* variants, the p.Val2050Leu variant in *ABCA4* was classified as likely benign, which makes it unlikely to be a modifier that worsens the phenotype.

The splice site variant c.828+3A>T was recurrently found in 57 index patients. This variant is predicted to result in aberrant splicing of the *PRPH2* messenger RNA, and consequently in the formation of a nonfunctional truncated protein (Shankar et al., 2015, 2016). The remaining 237 variants were reported in only a single or a few index patients, which clearly demonstrates the enormous allelic heterogeneity among patients with *PRPH2*-associated IRD.

4 | PATHOGENICITY ASSESSMENT OF ALL *PRPH2* VARIANTS

The pathogenicity of all *PRPH2* variants was assessed using the ACMG classification system, as described in the Materials and Methods section. According to our analysis, 107 variants were classified as pathogenic, 92 as likely pathogenic, one as benign, and two as likely benign (Figure S1B and Table S2). The remaining 50 variants were classified as variants of uncertain significance (Tables S2 and S3). Of the collective 199 (likely) pathogenic variants, 93 were missense, 85 were protein-truncating, 8 were splice-site, and 13 were in-frame amino acid insertions/deletions (Figure 1A and Table S2).

4.1 | Missense variants

The vast majority of *PRPH2* variants reported in IRD patients represent missense variants. In total, 137 missense variants have been reported, corresponding to 605 alleles (Figure S1B and Table 2). Before assessing the pathogenicity of these variants, in silico predictions were performed using PolyPhen2 (benign = 0.000–0.150; possibly damaging = 0.150–0.850; probably damaging = 0.850–1.000), SIFT (tolerated = 0.050–1.000; damaging = 0.000–0.050), and CADD (tolerated = <15; damaging = >15). To consider a variant as pathogenic (supporting evidence; PP3), two out of three in silico prediction programs should predict a damaging effect. When combining all information for the final ACMG pathogenicity assessment, of the 137 missense variants, 93 were classified as pathogenic/likely pathogenic (Figure 1B; Tables 3 and S2). One variant was classified as likely benign. The remaining 43 were considered as variants of uncertain significance (Tables S2 and S3), mainly due to lack of family history and segregation analysis, or because the variant did not co-segregate with the disease.

Next, we compared the position of (likely) pathogenic *PRPH2* missense variants relative to the protein structure of *PRPH2*, to identify regions that may be more prone to harbor disease-causing variants (Figure 1c). This analysis demonstrated that most missense variants (83 out of 93; Table 3; Figure 1c) are located within the D2 loop of the protein. It is hypothesized that the D2 loop is important for complex assembly with *ROM1* and/or dimerization with other *PRPH2* molecules, to both initiate and stabilize OS disc formation (Chakraborty et al., 2009, 2016; Goldberg et al., 1998). Furthermore, the D2 loop contains numerous highly conserved cysteine residues (Cys165, Cys166, Cys213, Cys214, Cys222, Cys250) that form disulfide bonds to maintain the structure of the loop and to regulate photoreceptor folding (Goldberg et al., 1998). For this reason, variants affecting amino acid residues within this loop will likely disrupt the structure or function of this loop, in turn interfering with *PRPH2*-*PRPH2* and/or *PRPH2*-*ROM1* interactions. Therefore, moderate evidence (PM1) was assigned if a variant was predicted to change the amino acid within this loop. Moderate evidence (PM1) was also assigned when missense variants were located in one of the transmembrane domains or C-terminus (Boon et al., 2008; Salinas et al., 2013).

For most of the missense variants, patient ethnicity was not always mentioned in the studied papers, which made it difficult to determine whether some variants might be specific for patients of a particular ancestry. Papers that did mention the ethnicity showed that most variants were reported in patients of Caucasian or Asian ancestry. Some variants seem to be specific for a certain ethnicity; for instance, the p.Cys250Gly variant was exclusively reported in patients with Asian ancestry, whereas p.Arg142Trp and p.Arg172Gln were only identified in patients with Caucasian ancestry. The p.Arg172Trp variant was almost exclusively reported in Caucasians but also reported once in a Japanese family (Nakazawa et al., 1995).

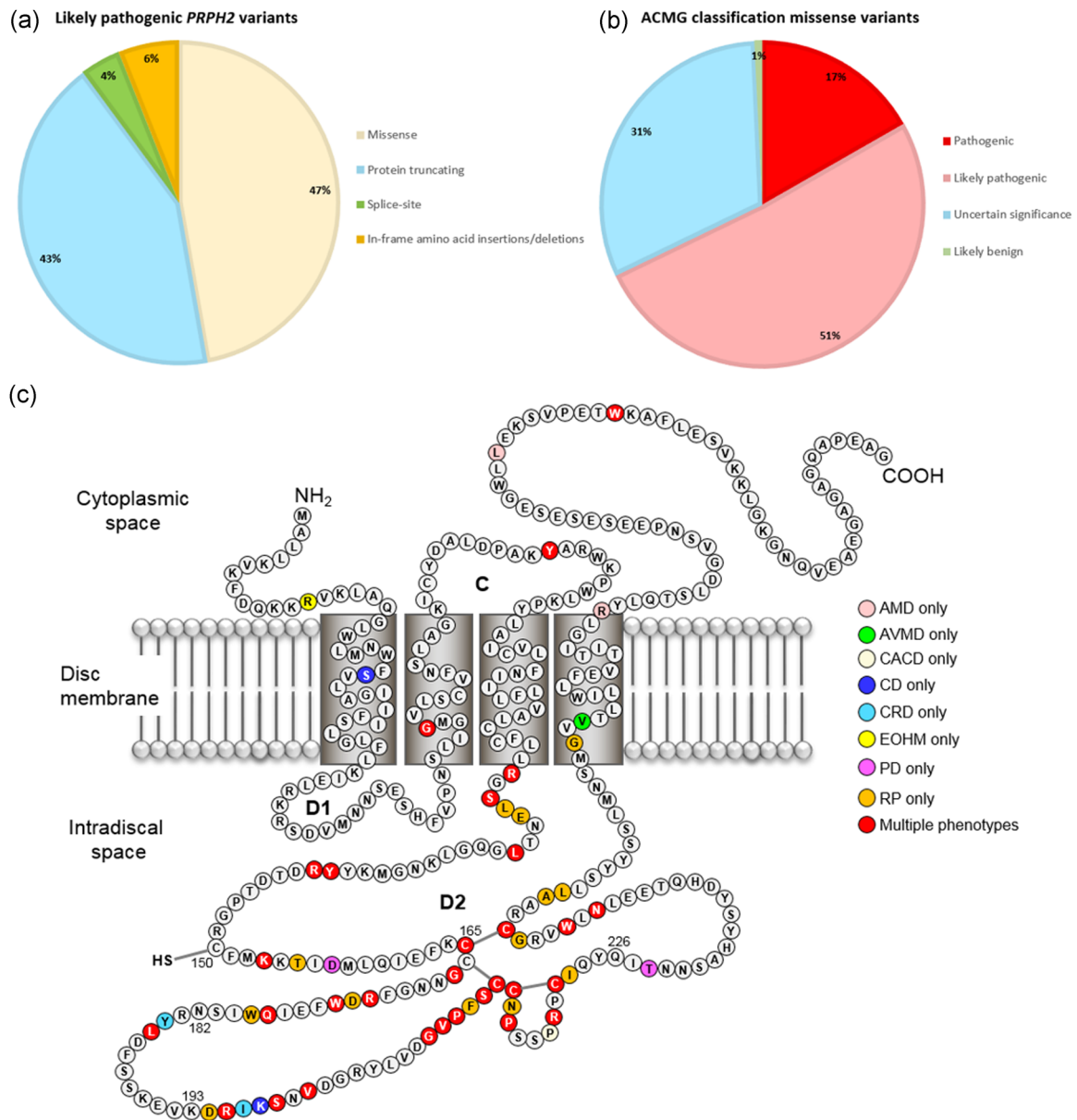


FIGURE 1 (a) Pie-chart showing the distribution of (likely) pathogenic *PRPH2* variants in IRD patients. (b) Pie-chart showing the ACMG pathogenicity assessment of missense variants. About two-thirds of the missense variants were classified as pathogenic or likely pathogenic. (c) Location likely pathogenic missense variants relative to the protein structure. The vast majority of the likely pathogenic missense variants are located in the D2 loop. AMD, age-related macular degeneration; AVMD, adult vitelliform macular dystrophy; CD, cone dystrophy; CRD, cone-rod dystrophy; EOHM, early-onset high myopia; IRD, inherited retinal disease; PD, pattern dystrophy; RP, retinitis pigmentosa

4.2 | Protein-truncating variants

In total, 180 out of 891 *PRPH2* alleles (20.2%) can be considered as protein-truncating, represented by 85 unique variants (Table 2). Of these, 55 were frameshift, 27 were nonsense, two were fail-to-start, and one was a deletion of exon 1. Most of these variants are considered rare, and only four of them were present in the gnomAD database. We classified a protein-truncating variant as damaging if the variant induces a premature stop codon before amino acid 331 since the p.Gln331* variant was reported to be pathogenic (Grover et al., 2002). Moreover, it

was determined that the valine residue at position 332 is critical for targeting *PRPH2* toward the OS of photoreceptor cells (Salinas et al., 2013). Based on this knowledge, we classified all protein-truncating variants as pathogenic or likely pathogenic (Table S2). One aspect to take into account is that the annotations of p.Tyr140ins 1bp, p.Ser217_dup16bp, and p.224ins-37bp, as described in the original articles, are not correct. We used Alamut to check whether we were able to identify the correct cDNA annotation, but this could not be deduced. For this reason, we kept the protein annotation used in the corresponding articles and put a question mark for the cDNA change annotation,

TABLE 3 Likely pathogenic and pathogenic missense variants

cDNA change	Protein change	Heterozygous	Compound heterozygous	Homozygous	Digenic	Trigenic	Protein domain
c.38G>A	p.Arg13Gln	2	0	0	0	0	N-terminus
c.80C>T	p.Ser27Phe	1	0	0	0	0	1st TMD
c.202G>C	p.Gly68Arg	2	0	0	0	0	2nd TMD
c.271T>A	p.Tyr91Asn	1	2	0	0	0	C-loop
c.271T>C	p.Tyr91His	1	0	0	0	0	C-loop
c.367C>T	p.Arg123Trp	2	0	0	0	0	D2-loop
c.374C>T	p.Ser125Leu	1	0	0	0	0	D2-loop
c.377T>C	p.Leu126Pro	3	0	0	0	0	D2-loop
c.377T>G	p.Leu126Arg	2	0	0	0	0	D2-loop
c.380A>G	p.Glu127Gly	2	0	0	0	0	D2-loop
c.389T>C	p.Leu130Pro	6	0	0	0	0	D2-loop
c.421T>C	p.Tyr141His	2	0	0	0	0	D2-loop
c.422A>G	p.Tyr141Cys	24	0	0	0	0	D2-loop
c.424C>T	p.Arg142Trp	93	0	2	0	0	D2-loop
c.425G>A	p.Arg142Gln	2	0	0	0	0	D2-loop
c.457A>G	p.Lys153Glu	1	0	0	0	0	D2-loop
c.458A>G	p.Lys153Arg	3	0	0	0	0	D2-loop
c.464C>T	p.Thr155Ile	0	0	0	1	0	D2-loop
c.469G>A	p.Asp157An	5	0	0	0	0	D2-loop
c.494G>T	p.Cys165Pe	1	0	0	0	0	D2-loop
c.494G>A	p.Cys165Tyr	4	0	0	0	0	D2-loop
c.493T>C	p.Cys165Ag	1	0	0	0	0	D2-loop
c.499G>A	p.Gly167Ser	10	0	0	0	0	D2-loop
c.500G>A	p.Gly167Asp	3	0	0	0	0	D2-loop
c.515G>A	p.Arg172Gln	13	0	0	0	0	D2-loop
c.514C>G	p.Arg172Gly	1	0	0	0	0	D2-loop
c.514C>T	p.Arg172Trp	58	0	0	1	1	D2-loop
c.518A>T	p.Asp173Val	4	0	0	0	0	D2-loop
c.520T>A	p.Trp174Arg	2	0	0	0	0	D2-loop
c.533A>G	p.Gln178Arg	2	0	0	0	0	D2-loop
c.536G>T	p.Trp179Leu	1	0	0	0	0	D2-loop
c.535T>C	p.Trp179Arg	4	0	0	0	0	D2-loop
c.537G>T	p.Trp179Cys	1	0	0	0	0	D2-loop
c.551A>C	p.Tyr184Ser	1	0	0	0	0	D2-loop
c.554T>C	p.Leu185Pro	6	0	2	4	0	D2-loop
c.582T>A	p.Asp194Gu	1	0	0	0	0	D2-loop
c.583C>G	p.Arg195Gly	1	0	0	0	0	D2-loop
c.584G>A	p.Arg195Gln	6	0	0	0	0	D2-loop
c.584G>T	p.Arg195Leu	12	0	0	0	0	D2-loop

(Continues)

TABLE 3 (Continued)

cDNA change	Protein change	Heterozygous	Compound heterozygous	Homozygous	Digenic	Trigenic	Protein domain
c.587T>A	p.Ile196Asn	1	0	0	0	0	D2-loop
c.589A>G	p.Lys197Glu	2	0	0	0	0	D2-loop
c.592A>C	p.Ser198Arg	1	0	0	0	0	D2-loop
c.594C>G	p.Ser198Arg	3	0	0	0	0	D2-loop
c.599T>A	p.Val200Glu	1	0	0	0	0	D2-loop
c.599T>G	p.Val200Gly	1	0	0	0	0	D2-loop
c.623G>A	p.Gly208Asp	12	0	1	0	0	D2-loop
c.625G>A	p.Val209Ile	2	0	0	0	0	D2-loop
c.625G>T	p.Val209Phe	1	0	0	0	0	D2-loop
c.626T>A	p.Val209Asp	2	0	0	0	0	D2-loop
c.628C>T	p.Pro210Ser	2	0	0	0	0	D2-loop
c.629C>G	p.Pro210Arg	20	0	1	0	0	D2-loop
c.629C>T	p.Pro210Leu	2	0	0	0	0	D2-loop
c.631T>C	p.Phe211Leu	6	0	0	0	0	D2-loop
c.634A>G	p.Ser212Gly	7	0	0	0	0	D2-loop
c.635G>C	p.Ser212Thr	4	0	0	0	0	D2-loop
c.636 T>A	p.Cys213Ser	2	0	0	0	0	D2-loop
c.637T>C	p.Cys213Arg	1	0	1	0	0	D2-loop
c.638G>T	p.Cys213Phe	1	0	0	0	0	D2-loop
c.638G>A	p.Cys213Tyr	6	0	0	0	0	D2-loop
c.639C>G	p.Cys213Trp	3	0	0	0	0	D2-loop
c.641G>A	p.Cys214Tyr	2	0	0	0	0	D2-loop
c.641G>C	p.Cys214Ser	1	0	0	0	0	D2-loop
c.643A>T	p.Asn215Tyr	1	0	0	0	0	D2-loop
c.646C>T	p.Pro216Ser	21	0	0	0	0	D2-loop
c.646C>G	p.Pro216Ala	1	0	0	0	0	D2-loop
c.647C>G	p.Pro216Arg	1	0	0	0	0	D2-loop
c.656C>G	p.Pro219Arg	2	0	0	0	0	D2-loop
c.658C>T	p.Arg220Trp	8	0	0	0	0	D2-loop
c.659G>A	p.Arg220Gln	3	0	1	0	0	D2-loop
c.659G>C	p.Arg220Pro	2	0	0	0	0	D2-loop
c.664T>C	p.Cys222Arg	2	0	0	0	0	D2-loop
c.665G>C	p.Cys222Ser	2	0	0	0	0	D2-loop
c.665G>A	p.Cys222Tyr	2	0	0	0	0	D2-loop
c.668T>A	p.Ile223Asn	1	0	0	0	0	D2-loop
c.683C>T	p.Thr228Ile	5	0	0	0	0	D2-loop
c.730A>C	p.Asn244His	1	0	0	0	0	D2-loop
c.732C>A	p.Asn244Lys	2	0	0	0	0	D2-loop
c.732C>G	p.Asn244Lys	1	0	0	0	0	D2-loop

TABLE 3 (Continued)

cDNA change	Protein change	Heterozygous	Compound heterozygous	Homozygous	Digenic	Trigenic	Protein domain
c.736T>C	p.Trp246Arg	2	0	0	0	0	D2-loop
c.738G>C	p.Trp246Cys	1	0	0	0	0	D2-loop
c.745G>A	p.Gly249Ser	1	0	0	0	0	D2-loop
c.748T>C	p.Cys250Arg	1	0	0	0	0	D2-loop
c.748T>G	p.Cys250Gly	3	1	0	1	0	D2-loop
c.748T>A	p.Cys250Ser	1	0	0	0	0	D2-loop
c.749G>A	p.Cys250Tyr	2	0	0	0	0	D2-loop
c.749G>T	p.Cys250Phe	1	0	0	0	0	D2-loop
c.758C>A	p.Ala253Glu	1	0	0	0	0	D2-loop
c.761T>A	p.Leu254Gln	5	0	1	0	0	D2-loop
c.797G>A	p.Gly266Asp	4	0	0	0	0	4th TMD
c.802G>A	p.Val268Ile	1	0	0	0	0	4th TMD
c.850C>T	p.Arg284Cys	1	0	0	0	0	C-terminus
c.923T>A	p.Leu308Gln	2	0	0	0	0	C-terminus
c.946T>G	p.Trp316Gly	5	0	0	0	0	C-terminus

Note: Novel missense variants are depicted in bold.

Abbreviations: 1st TMD, first transmembrane domain; 2nd TMD, second transmembrane domain; 4th TMD, fourth transmembrane domain; C-loop, cytoplasmic loop; D2-loop, second intradiscal loop.

although each is categorized as a protein-truncating variant. These variants are indicated in red in Table S2.

4.3 | Splice variants

Variants that were known to affect the canonical di-nucleotides of the splice acceptor (AG) or splice donor (GT) site were assigned to be pathogenic (PVS1). For all noncanonical splice site variants, four different splice prediction tools in Alamut were used to predict pathogenicity. A variant was classified as pathogenic when an increase or decrease of >10% in splice scores was observed for all four programs, as described previously (Messchaert et al., 2018). Based on this, together with the ACMG classification, almost all splice variants were classified as (likely) pathogenic (Table S2). There were two exceptions, namely c.581+4dupA and c.829-4C>G. For these variants, there was not enough robust evidence for their pathogenicity, and these were thus classified as variants of uncertain significance (Tables S2 and S3).

4.4 | In-frame amino acid insertions/deletions

Of the 15 in-frame insertions/deletions, 13 were classified as pathogenic or likely pathogenic. All in-frame amino acid insertions/deletions were assigned with PM4, indicating they might

affect normal protein length and function. In addition, none of these variants were present in gnomAD, indicating that these variants in fact might be disease-causing (Table S2). Two variants, namely p.Met67del and p.Met67_Gly68delinsArgHisArg, were classified as variants of uncertain significance (Table S3).

4.5 | Synonymous variants

Three synonymous variants were reported, namely p.Tyr101Tyr (c.303T>C), p.Tyr236Tyr (c.708T>C), and p.Glu335Glu (c.1005G>A). These variants were exclusively found in patients with macular dystrophy or pattern dystrophy. All variants were first classified as likely benign due to the fact that in silico prediction programs defined them as benign/tolerated. Moreover, no evidence for putative splice defects was predicted. However, the p.Tyr101Tyr (c.303T>C) and p.Glu335Glu (c.1005G>A) variants were not reported in the gnomAD database, indicating that they are not commonly found in control individuals (Table S2). For this reason, we classified these variants to be of uncertain significance (Tables S2 and S3). The p.Tyr236Tyr (c.708T>C) variant was reported in gnomAD, and statistical analysis showed that the variant was significantly enriched in the healthy control group when compared to index patients. Therefore, this variant was classified as likely benign (Table S2).

4.6 | Digenic and trigenic variants

In Section 3.2, it was shown that in a large German family segregating pattern dystrophy, p.Arg172Trp was present in patients that also carried a *ROM1* (p.Arg229His) and/or an *ABCA4* (p.Val2050Leu) variant (Poloschek et al., 2010). When looking at the severity of each individual's phenotype, it appeared that patients carrying an additional *ROM1* (digenic carriers) variant were more severely affected compared to patients that only carried the *PRPH2* p.Arg172Trp variant. The phenotype was even more severe in patients carrying *PRPH2*, *ROM1*, and *ABCA4* variants (trigenic carriers). The authors hypothesized that the additional *ROM1* and *ABCA4* variants function as genetic modifiers that worsen the pattern dystrophy phenotype. However, the p.Val2050Leu variant in *ABCA4* on itself was classified as likely benign, which makes it unlikely to be a modifier worsening the disease phenotype.

The p.Leu185Pro variant in *PRPH2* was also reported together with *ROM1* variants (p.Gly80Gly, p.Gly113Glu, or p.Leu114Leu) in four large digenic RP families (Dryja et al., 1997; Kajiwara et al., 1994). In these families, individuals only carrying the p.Leu185Pro variant seemed to remain unaffected by disease, while individuals that also carried one of the *ROM1* variants showed clear phenotypic characteristics of RP. The p.Leu270del variant in *PRPH2* was found in combination with the p.Met318Alafs*17 variant in *ROM1* in another digenic RP patient (Dryja et al., 1997). Recently, *PRPH2* p.Arg46* in combination with the complex p.([Leu2027-Phe];(Gly1977Ser)] *ABCA4* variant, was identified in a patient with cone-rod dystrophy (Coco-Martin et al., 2020). Finally, the same group also reported a patient with pattern dystrophy carrying both p.Lys154del in *PRPH2* and p.Arg2030Gln in *ABCA4* (Coco-Martin et al., 2020). These findings indicate that besides RP, also cone-rod and pattern dystrophy might be caused by digenic variants (i.e. one in *PRPH2* and one in another gene). However, mutations in IRD associated genes are relatively frequent. Nishiguchi and Rivolta screened high-quality genome sequences of control individuals from various ancestries and estimated that ~1 in 4–5 individuals from the general population carry variants that are associated with IRD (Nishiguchi & Rivolta, 2012). Furthermore, Hanany and colleagues created a database containing 276,921 sequence variants that were identified in 187 autosomal recessive IRD genes and found that 2.7 billion people worldwide (36% of the population) are being healthy carriers of at least one IRD variant (Hanany et al., 2020). Thus, patients may carry variants in multiple IRD genes without any clinical consequences. For this reason, caution is warranted, and studies of large cohorts are required to determine if the disease in a patient is truly inherited in a di- or trigenic fashion.

4.7 | Reclassified variants

The evaluation of the pathogenicity for some variants resulted in a different outcome when compared to the original publication. This is partly due to the fact that more specific variant classification tools

have become available. For example, the p.Tyr101Tyr (c.303T>C) and p.Glu335Glu (c.1005G>A) variants were previously classified to be benign by Dryja et al. (1997), mainly since they do not change the amino acid sequence, and thus are predicted not to have a deleterious effect on the final protein. However, we have classified these variants as being of uncertain significance since they were not present in normal controls reported in gnomAD, and thus in fact may be disease-causing. Experimental assays are needed to truly determine the potential causality of these variants. Another example is the c.-11A>C variant, which is located in the 5'-UTR region. This variant was classified as likely pathogenic because it was not present in single-nucleotide polymorphism databases, and was not found in 92 controls (Boon, Klevering, et al., 2007). However, we have classified the variant as benign since when comparing the allele frequency in gnomAD to the allele frequency in the reported index patients, the variant was found to be enriched in the control population (gnomAD).

There were also some variants that were considered to be of uncertain significance in initial publications but were classified as (likely) pathogenic following our classification. For instance, the p.Gly167Ser variant (Meunier et al., 2011; Strom et al., 2012) is located in the D2 loop (mutational hotspot; PM1), was absent in controls (gnomAD; PM2), was predicted to be damaging by all three in silico prediction programs (PP3; Table S2), and co-segregated with disease (PP1). Another mis-classified variant was p.Gly68Arg, originally considered to be of uncertain significance (Dryja et al., 1997), and found in patients that were suspected to have digenic RP. However, analysis of the segregation of *ROM1* alleles was uninformative. For this reason, the authors could not determine whether the p.Gly68Arg defect is pathogenic or that it represents a rare benign variant. Upon our pathogenicity classification, the variant was classified as likely pathogenic since it was significantly enriched in the patient population compared to controls (gnomAD; PS4), was located in the first TMD (PM1), and was predicted to be damaging by all three in silico prediction programs (PP3; Table S2). These findings demonstrate that pathogenicity classification tools have improved, enabling reclassification of certain variants that were reported many years ago.

5 | GENOTYPE-PHENOTYPE CORRELATIONS

To define genotype-phenotype correlations, we collected information about disease phenotype, age at onset, and, when available, segregation data. However, most of the *PRPH2* variants show high phenotypic variability, both between different families and within the same family, which made it difficult to draw proper and well-defined conclusions regarding the relationship between *PRPH2* variants and disease phenotype. For instance, in an Italian family, the p.Cys165Arg variant resulted in clinically different phenotypes (fundus flavimaculatus or butterfly-shaped pattern dystrophy) within the same family (Simonelli et al., 2007). The mechanism behind this intrafamilial phenotypic variability remains to be elucidated but the authors

suggested the following explanation; since *PRPH2* interacts with other adhesion proteins, the inheritance of genes encoding such proteins may—to some extent—explain the enormous phenotypic variability observed within families. A family described by Daftarian et al. carries both the p.Gln239* and the p.Ile32Val variant (Daftarian et al., 2019). The proband carried both variants in a homozygous state, indicating that these two variants are on the same allele, which resulted in the more severe phenotype LCA. Family members carrying the variants in a heterozygous state developed much milder phenotypes with a later age of onset. The authors concluded that homozygous variants result in more severe phenotypes compared to heterozygous *PRPH2* variants. A family with even higher phenotypic variability is a large Dutch family (family E) described by Boon et al. Here, similarly-aged family members showed phenotypes ranging from retinitis pigmentosa, pseudo-Stargardt pattern dystrophy, to only mild foveal pigmentary changes, despite carrying the same frameshift mutation (p.Gly148Alafs*5) (Boon, van Schooneveld, et al., 2007). The broad spectrum of phenotypical variability associated with *PRPH2* variants is further highlighted by a reduction in penetrance described in several papers. For example, Michaelides et al. described a five-generation family in which two individuals carrying the p.Arg172Trp mutation, a mother (49 years old) and her daughter (24 years old), had an entirely normal phenotype upon detailed testing (Michaelides et al., 2005). As previously mentioned, Boon et al. also described a normal phenotype in patients carrying the p.Arg142Trp mutation, even up until the age of 64 years (Boon et al., 2009), illustrating the reduced penetrance for (at least some) *PRPH2* mutations. Another point to address is that we did not observe obvious indications that individuals with protein-truncating variants had a more severe phenotype than, for instance, individuals carrying missense variants or in-frame amino acid insertions/deletions. Finally, when looking at the location of likely pathogenic *PRPH2* variants, it was striking that these variants seem to cluster in the D2 loop (Figure 1c), suggesting that this loop is a mutational hot spot for amino acid changes associated with *PRPH2*-associated IRD. Taken together, additional genetic data from IRD patients would be of great help to determine genotype-phenotype correlations and to study the often observed intra- and inter-familial phenotypic variability.

6 | DISCUSSION

6.1 | Improvement of in silico analysis

Before classifying the pathogenicity of *PRPH2* missense variants according to ACMG, in silico predictions were performed using Polyphen-2, SIFT, and CADD. Polyphen2 is a software program and is accessible via a Web server. The program predicts the possible effects of non-synonymous single-nucleotide variants (SNVs) on the stability and function of human proteins. It uses both structural and comparative evolutionary considerations. These properties are then combined to estimate the probability of a particular missense variant being damaging to the protein of interest (Adzhubei et al., 2013).

SIFT uses datasets of functionally related protein sequences that are obtained via a protein database. The algorithm scans each position in the sequence of interest, after which the probabilities for all possible 20 amino acids at that position are calculated, resulting in one final SIFT score (Kumar et al., 2009). The CADD in silico prediction software is also available via a webserver. The program integrates multiple lineages of data, including genomic features within the surrounding sequence of interest, gene model annotations, evolutionary data from multiple species, epigenetic measurements, as well as functional predictions. In contrast to many other in silico prediction programs, CADD is not limited to the number of genomic variants of which either a pathogenic or benign status is already known (Rentzsch et al., 2019). Instead, CADD bases its score on less biased and much larger data sets and thus might have an advantage over Polyphen2 and SIFT.

Some missense variants, including p.Ala2Ser, p.Lys15Arg, p.Ser27Phe, p.Arg123Trp, p.Ser125Leu, p.Glu127Gly, p.Arg142Gln, p.Thr155Ile, p.Arg172Gln, p.Asp186Asn, p.Asp194Glu, p.Ile196Phe, p.Lys197Glu, p.Asn199Asp, p.Gly208Asp, p.Ser212Thr, p.Pro216Leu, p.Thr228Ile, p.Glu242Gly, p.Ala252Pro, p.Leu261Phe, p.Trp316Gly, and p.Ala337Thr, gave contradictory outcomes upon in silico pathogenicity predictions. Other variants, including p.Ile32Val, p.Ala85Thr, p.Ala116Ser, p.Met152Val, p.Lys154Gln, p.Ile161Met, p.Ser217Gly, p.Leu261Phe, p.Val268Ile, p.Pro313Ser, and p.Pro313Leu were predicted to be benign. While different algorithms may rely on the same type of data to predict pathogenicity, it is known that for some genes, similar algorithms can have a significantly different outcome (Richards et al., 2015). Only when at least two out of three of the in silico programs used give the same prediction output, the evidence can be counted as supporting (PP3) (Richards et al., 2015). Based on this, the in silico predictions for the p.Lys15Arg, p.Ile32Val, p.Ala85Thr, p.Ala116Ser, p.Glu127Gly, p.Met152Val, p.Lys154Gln, p.Ile161Met, p.Asp186Asn, p.Asp194Glu, p.Ile196Phe, p.Pro216Leu, p.Ser217Gly, p.Thr228Ile, p.Leu261Phe, p.Val268Ile, p.Pro313Ser, and p.Pro313Leu, p.Trp316Gly, and p.Ala337Thr variants were not considered to be of supporting evidence (PP3). It would therefore be very helpful if there were more experimental data available that could show whether these variants have a deleterious effect on normal protein morphology and/or function. This will be explained in more detail in the next section.

6.2 | Experimental assessment of variants of unknown significance

Approximately one-third of the missense variants were classified as variants of unknown significance (Figure 1b). This was due to the fact that these variants did not robustly meet important ACMG pathogenicity criteria. Furthermore, two splice-site, two in-frame amino acid insertions/deletions, two synonymous variants, and one 3'-UTR variant also needed to be classified as variants of uncertain significance for the same reasons as for the missense variants.

Experimental models would be of great help to classify such variants. However, experimental data are extremely limited. This is mainly due to the fact that *PRPH2* expression is highly retina-specific. This makes it difficult to use widely used immortalized lymphoblastoid, skin fibroblast, or blood cell lines. However, depending on the type and location of each variant, this issue may be circumvented. For instance, when studying missense variants, one could clone the coding region of *PRPH2* (~1.1 kb) into specific expression vectors to create a *PRPH2* wildtype vector. Next, site-directed mutagenesis can be applied to insert the desired mutation. Wildtype and mutant vectors can be transfected into widely used cell lines to study, for example, expression patterns (western blots) or interacting proteins (yeast-two-hybrid, co-immunoprecipitation), and compare mutant and wild-type conditions. These vectors can even be administered to neonatal mice to study *in vivo* localization of wildtype or mutant *PRPH2*. Chakraborty and colleagues created vectors containing wild-type *PRPH2* or the *PRPH2* p.Cys213Tyr variant and electroporated these into neonatal mice. Four weeks after injection, retinas were collected and it was seen that the p.Cys213Tyr construct mislocalized to the inner segments and perinuclear region (Chakraborty et al., 2020). Another research group showed that constructs containing the p.Pro210Leu and p.Cys214Ser missense variants mislocalized to the inner segments upon administration to wildtype mice (Becirovic et al., 2016). Interestingly, results differed between rods and cones, indicating that *PRPH2* might have a different function in the two different photoreceptor cell types.

Second, when one wants to study splice site variants, minigene or midigene splice assays can be used, as described previously (Sangermano et al., 2018). Minigenes or midigenes represent plasmids in which the desired *PRPH2* splice site variant can be cloned between two exons of interest. The plasmids are transfected into, for example, HEK293T cells, and splice effects can be analyzed using reverse-transcription polymerase chain reaction (RT-PCR) to determine whether there are differences between RNAs transcribed from wild-type and mutant minigenes. Becirovic and colleagues constructed *PRPH2* wild-type minigenes as well as minigenes containing the IRD-associated p.Arg195Leu, p.Ser198Arg, p.Val209Ile, p.Pro210Leu, p.Ser212Thr, p.Cys214Ser, p.Arg220Trp, p.Arg220Gln, p.Trp246Arg, and p.Gly249Ser missense variants (Becirovic et al., 2016). RT-PCR analysis of *PRPH2* transcripts in murine retinas transduced with the wild-type or the mutant *PRPH2* minigene identified three different *PRPH2* splice isoforms in rods and cones. These splice isoforms consisted of the unspliced transcript, a transcript in which intron 1 was retained, and the correctly spliced *PRPH2*. The p.Gly249Ser variant created a new splice donor site, resulting in aberrant splicing of the protein. This suggests that also point mutations in coding regions might affect splicing (Becirovic et al., 2016).

Finally, for studying all types of variants, one could ideally make use of induced pluripotent stem cell (iPSC) technology. In short, somatic cells are extracted from *PRPH2*-associated IRD patients or from healthy controls, which can then be reprogrammed toward iPSCs. Next, these iPSCs can be differentiated toward retina-like cells thus carrying the variant of interest (Giacalone et al., 2016; Öner, 2018), after which a variety of

functional studies can be performed, including expression (western blot), localization (immunohistochemistry), and morphological (microscopy) studies. A drawback of this approach is that it is very labor- and time-consuming, as well as expensive.

Taken together, these experimental studies are crucial to improve the classification of *PRPH2* variants and thereby aid in the molecular diagnostics of IRDs.

6.3 | Phenotypic variability and genotype-phenotype correlations

As detailed in the introduction and genotype-phenotype correlation section, there is significant variability in clinical phenotypes, with regard to age at onset, severity, and incomplete penetrance. Some variants show a clear genotype-phenotype correlation like p.Leu126Pro, p.Leu126Arg, p.Glu127Gly, p.Gly137Asp, p.Arg142QGln, p.Met152Val, p.Cys165Tyr, p.Asp173Val, p.Trp179Arg, p.Ser198Arg, p.Pro210Ser, p.Phe211Leu, and p.Gly266Asp, and that all seem to be associated with the development of RP only (Anasagasti et al., 2013; Bareil et al., 2000; de Sousa Dias et al., 2015; Dryja et al., 1997; Gao et al., 2019; Grüning et al., 1994; Jin et al., 2008; Kemp et al., 1994; Manes et al., 2015; Reeves et al., 2020; Renner et al., 2009; Sohocki et al., 2001; Souied et al., 1998; Stone et al., 2017; Sullivan et al., 2013; Van Cauwenbergh et al., 2017; Yoon et al., 2015). Other examples are p.Asp157Asn and p.Cys213Trp, which are exclusively associated with PD (Boon, van Schooneveld, et al., 2007; Jacobson et al., 1996; Reeves et al., 2020). Thus, these disorders can best be described as allelic disorders due to the fact that there is a clear variant-phenotype relationship. However, for most of the variants detailed in this study, there appears to be a high phenotypic variability between different families, as well as between members within the same family. A clear example of this is explained by the c.828+3 A>T variant described by Shankar and colleagues. This particular variant results in the development of multiple distinct IRD phenotypes, including RP, CRD, PD, and macular dystrophy (Shankar et al., 2015, 2016). The multiple phenotypes that are seen in these families can best be referred to as variable expressivity as was also concluded by the authors of the respective paper. Another example also showing variable expressivity is p.Arg142Trp. This variant is associated with multiple IRD phenotypes that, in some cases, also show incomplete penetrance.

6.3.1 | Insights from the molecular pathophysiology of the *PRPH2* p.Arg172Trp variant

When looking at the subtype of IRD in patients carrying the p.Arg172Trp variant, it becomes apparent that these subjects have a relatively young mean age of onset, and an overall cone-dominant phenotype (Reeves et al., 2020; Wells et al., 1993; Wroblewski et al., 1994a). Most of the patients display a significant decrease in cone function, while rod function seems to remain unaffected (Conley et al., 2007). In a rare number of cases, however, usually observed in later stages of the disease, rods become affected as well. Expression of the human-specific p.Arg172Trp

variant in mice resulted in the rescue of the rod but not cone phenotypes. This finding strongly indicates that multiple variant-specific mechanisms exist (Ding et al., 2004).

Co-expression of the p.Arg172Trp mutation and a full complement of wildtype Prph2 initially resulted in normal rod function and structure (Conley et al., 2014). For cones, however, something different was observed. The retina of mice carrying the Arg172Trp variant exhibited a significant loss in the total number of blue and green cones, as well as a decrease in photopic ERG responses. Furthermore, defects in Prph2-Rom1 complex formation were observed (Conley et al., 2014). In another study, the p.Arg172Trp variant was expressed on a rod dominant (Prph2^{+/-}) background and led to a late-onset reduction in the number of rod cells as well as scotopic responses. The authors speculated that this was highly likely due to the haploinsufficiency of Prph2 (Conley et al., 2014). Expression of the Arg172Trp mutation in the cone-dominant (Nrl^{-/-}) retina resulted in a functional decline of the photopic responses, as well as disrupted cone OS structures, due to the formation of abnormal high molecular weight Prph2-Rom1 aggregates (Conley et al., 2014).

Based on the aforementioned, we can conclude that the p.Arg172Trp variant is most frequently associated with cone-dominant phenotypes and that multiple variant-specific mechanisms underlying the development of IRD exist. Together, the exact mechanisms by which PRPH2 mutations can lead to one or more distinct clinical subtypes of IRD for a large part remain elusive.

6.4 | Development of therapeutic strategies for PRPH2-associated IRD

Proper classification of possible disease-causing variants is of great importance, not only to determine the true causal genetic defect in IRD patients but also for developing therapeutic strategies. PRPH2 has been a target for gene therapy for over two decades now, mainly due to the disease burden associated with variants in this gene, as well as the availability of several extremely well-characterized animal models mimicking important phenotypic characteristics of patients with PRPH2-associated IRD (Conley & Naash, 2014). Currently, at least three therapeutic strategies can be distinguished: (1) gene replacement therapy; (2) gene knockdown therapy; and (3) delivery of neurotrophic factors. To develop such therapeutic strategies, it is important to not only know the genetic defect but also the underlying pathophysiological mechanism (e.g., dominant-negative vs. haploinsufficiency).

6.4.1 | Gene replacement therapy

The first proof-of-principle studies regarding gene replacement therapy attempted to correct IRD phenotypes in the Prph2^{-/-} and Prph2^{+/-} mice. For this purpose, a wild-type Prph2 transgene was delivered, and the results were highly promising since in Prph2^{-/-} mice, the structure of the OS of rod photoreceptor cells was

preserved (Travis et al., 1992). Furthermore, in Prph2^{+/-} mice, the expression of a wildtype Prph2 transgene rescued rod and cone OS structure and function (Nour et al., 2004). In mice harboring the recurrent p.Arg172Trp variant, which is considered a dominant-negative variant, expression of wildtype PRPH2 also caused structural and functional improvements (Conley et al., 2007; Nour et al., 2008).

6.4.2 | Gene knockdown therapy

As some mutations in PRPH2-associated IRD are believed to act in a dominant-negative manner, such as p.Arg172Trp (Conley et al., 2007), and the fact that gene replacement did not completely correct the dominant phenotype in mice carrying this specific variant, alternative approaches are needed to eliminate the mutant allele. One such approach is the gene knockdown approach. For PRPH2, the usage of short hairpin RNA (shRNA) to knock down the mutant allele seems to be the most promising. For example, an shRNA that was shown to knock down Prph2 in vitro was packaged into an rAAV vector. Upon injection into the subretinal space of wild-type mice, wild-type Prph2 levels were reduced by 75% (Peters-Silva et al., 2012). Next, they created a vector containing wildtype Prph2 that was proven to be resistant to the aforementioned shRNA. Co-delivery of these vectors resulted in the rescue of functional defects caused by the shRNA knockdown and partial recovery of total Prph2 levels (Peters-Silva et al., 2012). Similarly, using electroporation instead of a virus, coinjection of both an shRNA vector that is able to knock down both wildtype and mutant Prph2 and an shRNA-resistant copy of wild-type Prph2 resulted in degradation of endogenous Prph2 and stabilized expression of the exogenously delivered Prph2 in mouse retinal explants (Palfi et al., 2006). Although the efficacy of this kind of therapy has not yet been evaluated in IRD disease models, these studies show that allele-independent knockdown in combination with gene supplementation represents a potential therapy to counteract genetic defects in PRPH2-associated diseases.

6.4.3 | Delivery of neurotrophic factors

Unlike gene replacement and gene knockdown therapies, also more general therapeutic strategies are considered, for example, the delivery of neurotrophic factors. An advantage of such a strategy is that it can be applied to multiple genetic subtypes of IRD. The first proof-of-principle came from a mouse study in which ciliary neurotrophic factor (CNTF) was injected into the intravitreal space of Prph2^{-/-} mice (Cayouette et al., 1998), and lead to an improvement in OS structure. However, at the functional level, only a small improvement in rods and no improvement in cones was observed. In a similar study performed by another group, CNTF was delivered to p.Pro216Leu Prph2^{+/-} mice. These mice showed improved OS structure, but the authors also observed dose-dependent abnormalities in photoreceptor nuclei as well as a decrease in both rod and cone function

(Bok et al., 2002). Follow-up studies revealed that CNTF alters retinal signaling pathways. Furthermore, they observed a downregulation of critical phototransduction genes, such as cone opsins (Rhee et al., 2007). Although these adverse findings have led to the preclusion to use CNTF to treat *PRPH2*-associated IRD, other neurotrophic factors were investigated. For example, lentiviruses carrying either fibroblast growth factor-2 (FGF-2) or human pigment epithelial-derived factor (PEDF) were reported to significantly improve both scotopic a- and b-waves in *Prph2*^{-/-} mice. However, these agents did not restore photoreceptor survival (Miyazaki et al., 2008). Other nontraditional neurotrophic factors, such as hormones, also have been shown to result in neuroprotection. Administration of some of these agents, including erythropoietin (hormone) (Rex et al., 2004, 2009), and nilvadipine (calcium channel blocker) (Takeuchi et al., 2008), significantly improved photoreceptor function in *Prph2*^{-/-} and *Prph2*^{+/-} mouse models. However, more studies are required to determine both the safety and efficacy of such particular therapeutic approaches.

7 | CONCLUDING REMARKS

Taken together, in this study, we describe 245 reported and seven novel *PRPH2* variants identified in 891 alleles in 867 index cases and uploaded these to the LOVD and ClinVar database for *PRPH2*. This study thereby provides an important step toward a complete overview of all *PRPH2* variants in a single database. Continuous addition of genetic data from newly identified patients with *PRPH2* variants is of great importance for more robust classification of pathogenic variants. Furthermore, additional data regarding phenotypes might aid the identification of genotype-phenotype correlations. Our analysis resulted in the *in silico* classification of all the 245 reported, as well as the seven novel identified *PRPH2* variants. More importantly, our study illustrates the need for experimental assays to identify the true causality of the many *PRPH2* variants that are now still assigned to be variants of uncertain significance. This will help to improve molecular diagnostics and, in the long-term, hopefully also support the development of therapeutic strategies for patients with *PRPH2*-associated IRD.

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CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

WEB RESOURCES

<https://gnomad.broadinstitute.org/>
<http://genetics.bwh.harvard.edu/pph2/>
<http://cadd.gs.washington.edu/home>
<https://sift.bii.a-star.edu.sg/>
<https://databases.lovd.nl/shared/genes/PRPH2>
[https://www.ncbi.nlm.nih.gov/clinvar/?term=PRPH2\[all\]](https://www.ncbi.nlm.nih.gov/clinvar/?term=PRPH2[all])

DATA AVAILABILITY STATEMENT

All variant data described in this study are freely available via the LOVD and ClinVar database for *PRPH2* (<https://databases.lovd.nl/shared/variants/PRPH2>; PRPH2[all] - ClinVar - NCBI (nih.gov)). Other data that support the findings detailed in this study are available from the corresponding author, upon reasonable request.

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SUPPORTING INFORMATION

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