CRB1-associated Retinal Dystrophies: Genetics, Clinical Characteristics and Natural History.

Running head: CRB1 Retinal Dystrophy Natural History

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Abstract

**Purpose:** To analyse the clinical characteristics, natural history, and genetics of CRB1-associated retinal dystrophies.

**Design:** Multicenter international retrospective cohort study.

**Methods:** Review of clinical notes, ophthalmic images, and genetic testing results of 104 patients (91 probands) with disease-causing CRB1 variants. Macular optical coherence tomography (OCT) parameters, visual function, fundus characteristics, and associations between variables were our main outcome measures.

**Results:** The mean age of the cohort at the first visit was 19.8 ± 16.1 (median 15) years of age, with a mean follow-up of 9.6 ± 10 years. Based on history, imaging, and clinical examination, 26 individuals were diagnosed with retinitis pigmentosa (RP, 26%), 54 with early-onset severe retinal dystrophy/Leber Congenital Amaurosis (EOSRD/LCA, 51%), and 24 with macular dystrophy (MD, 23%). Severe visual impairment was most frequent after 40 years of age for patients with RP and after 20 years of age for EOSRD/LCA.

Longitudinal analysis revealed a significant difference between baseline and follow up best corrected visual acuity in the three sub-cohorts. Macular thickness decreased in most patients with EOSRD/LCA and MD, whereas the majority of patients with RP had increased perifoveal thickness.

**Conclusions:** A subset of individuals with CRB1 variants present with mild, adult-onset RP. EOSRD/LCA phenotype was significantly associated with null variants, and 167_169 deletion was exclusively present in the MD cohort. The poor OCT lamination may have a degenerative component, as well as being congenital. Disease symmetry and reasonable window for intervention highlight CRB1 retinal dystrophies as a promising target for trials of novel therapeutics.
INTRODUCTION

Biallelic disease-causing variants in Crumbs homolog 1 (CRB1, MIM# 604210) have been associated with a wide and complex range of phenotypes. The most commonly reported is Leber congenital amaurosis (LCA)/early onset severe retinal dystrophy (EOSRD), where CRB1 accounts for around 10% of all cases.\(^1\) This is followed in frequency by retinitis pigmentosa (RP),\(^2,3\) in which CRB1 represents up to 6.5%.\(^4\) Other phenotypes include cone-rod (CRD),\(^5,6\) macular dystrophy (MD),\(^7,8\) foveal retinoschisis,\(^9\) and fenestrated slit maculopathy.\(^10\)

One of the distinctive features of CRB1-retinopathy can be the presence of nummular pigmented deposits,\(^11\) admixed with small yellow-white dots.\(^12\) Also, CRB1 is commonly associated with preserved para-arteriolar retinal pigment epithelium (PPRPE),\(^2,13\) abnormally laminated and thickened retina,\(^14\) and peripheral exudative retinal telangiectasia (Coats-like vasculopathy)\(^15,16\) - which can ultimately lead to retinal detachment and neovascular glaucoma.\(^5\) Non-retinal features have also been linked to this gene such as nanophthalmos,\(^17,18\) hyperopia,\(^13,19\) narrow anterior chamber,\(^20\) and optic disc drusen.\(^18\)

CRB1 was first identified in several unrelated individuals with RP and PPRPE.\(^21,22\) CRB1 encodes a transmembrane protein with multiple epidermal growth factor-like and laminin A globular-like domains;\(^15\) and is believed to have a role in retinal development as well as in long-term retinal integrity. Its primary function is in the maintenance of the zonula adherens junctions between photoreceptors, Müller glial cells and the external limiting membrane (ELM).\(^23-25\) It also has essential roles in epithelial cell polarity and in the scaffolding complex, in vascular integrity, and is key to the preservation of an organized photoreceptor layer.\(^26\)

CRB1 is frequently reported as one of the most common causative genes for LCA/EOSRD.\(^27-29\) This has driven increasing efforts to develop animal models and treatments.\(^30,31\) As the preclinical work moves forward, it becomes imperative to understand the natural history of the disease. In this retrospective international study, we undertake deep phenotyping of the largest CRB1 cohort to date, report disease natural history and explore potential endpoints for future interventional clinical trials.
METHODS

The study protocol adhered to the tenets of the Declaration of Helsinki and was approved by the ethics committees of the participating institutions.

Patient Selection and Genetics

The inclusion criterion for the current study was to have molecularly confirmed CRB1-associated retinopathy. This was defined as patients with an inherited retinal dystrophy (IRD) harbouring 2 or more disease-causing CRB1 variants. The patients were identified by reviewing the genetics database of Moorfields Eye Hospital (London, UK) and St. John of Jerusalem Eye Hospital group (Jerusalem), and their records were subsequently studied.

Genetic testing was performed with various available methods, such as direct Sanger sequencing, next-generation sequencing (NGS)-based retinal dystrophy gene panels, whole exome sequencing (WES), and whole genome sequencing (WGS). In silico molecular genetic analysis was performed for all detected CRB1 variants (transcript reference: NM_201253.3: ENST00000367400.3) and the detailed description is provided in the supplemental material (Supplementary Methods, available at AJO.com). Pathogenicity of each variant was classified mainly according to the guidelines of the American College of Medical Genetics and Genomics (ACMG). The cut off value of allele frequency to apply PM2 (absence or very rare in the general population database) was 0.001. For the purpose of this study, an additional specification (Likely pathogenic: 1 Moderate (PM1-PM6) AND 3 Supporting (PP1-PP5)) to determine the verdict assessment results was applied to the ACMG classification.

Clinical Assessment and Retinal Imaging

All participants were seen by specialists in IRDs at referral sites. Clinical notes were reviewed, including family, medical and ophthalmic history, best corrected visual acuity (BCVA), refraction, slit-lamp biomicroscopy findings, and fundoscopy. BCVA was
converted to logarithmic minimum angle of resolution (LogMAR) for statistical analysis. Count fingers vision was given a value of LogMAR 1.98 and hand motion, LogMAR 2.28, light perception and no light perception were LogMAR 2.7 and 3, respectively.\textsuperscript{34,35} Patients were categorised using the World Health Organization (WHO) visual impairment criteria, that defines no or mild visual impairment as BCVA < 0.48 (6/18, 20/60), moderate impairment as BCVA > 0.48 and ≤ 1.0 (6/60, 20/200), severe as BCVA >1.0 and ≤ 1.3 (3/60, 20/400), and blindness as BCVA > 1.3. Records of visual field were limited within our cohort; therefore, we only took into consideration central vision based on BCVA to classify patients. 'Low vision' corresponds to patients with moderate and severe impairment. Asymmetric BCVA was defined as a difference greater than ≥0.3 LogMAR (equivalent to 15 ETDRS letters) between eyes. Refraction was undertaken by an optometrist for both adults and children, and spherical equivalent was calculated for refractive error.

When available, we also assessed additional testing such as color and autofluorescence retinal imaging, near-infrared reflectance, and OCT (details in Supplementary Methods). Fovea-centred macular volume scans were performed in a 6 mm\textsuperscript{2} area that included the standard 1, 3, and 6 mm grid template from the ETDRS. Inner limiting membrane and Bruch’s membrane were automatically segmented by the manufacturer software (Hefex version 1.9.14.0; Heidelberg Engineering) or adjusted manually as needed by a trained ophthalmologist (M.D.V.). Macular OCTs were divided into three categories regarding their qualitative features and associations subsequently explored: group 1, characterized by normal lamination; group 2, where the retinal layers are generally discernible but appear ill-defined; and group 3, defined by a disorganised retina with coalescent layers (particularly within the inner retina) and a degree of increased reflectivity of the nuclear layers (Figure 1).\textsuperscript{36,37} To allow direct comparison between normal and abnormal retinal architecture, we also compared normal (group 1) and abnormal OCT lamination (groups 2 and 3). Patients with macular cysts, edema, and/or only line scans due to poor fixation were excluded from quantitative assessments (details in Supplementary Methods). Normative data regarding OCT thickness in the general population was extracted from Grover et al.,\textsuperscript{38} and volume parameters were taken from Roshandel et al.\textsuperscript{10} Both eyes from each patient were analysed.
Electrophysiological assessment

Pattern and full-field electroretinogram (PERG; ff-ERG) testing was performed in a subset of patients, incorporating the standards of the International Society for Clinical Electrophysiology of Vision (ISCEV) and included additional dark-adapted (DA) red flash ERGs. The PERG and ff-ERG were performed using gold foil electrodes, except in 9 young children who underwent testing with lower eyelid skin electrodes using a shorter ERG protocol. The quantitative ERG data analysis was limited to recordings from Moorfields Eye Hospital, to optimise consistency of methods and to minimise variability due to different types of recording electrode. The patient ERG data were compared with those from a control group of healthy subjects (age range: 10-79 years). Further details can be found in the Supplementary Methods.

Statistics

Statistical analysis was carried out with GraphPad Prism 8.0.2 (GraphPad Software, San Diego, CA, USA). The threshold of significance for all statistical tests was set at \( p < 0.05 \). Linear regressions and t-test were used for parametric variables’ assessment. Welch’s t-test variation was employed when the sample sizes were significantly different. Chi-square was undertaken to assess possible association between two categorical variables.

RESULTS

Demographics, phenotypic category, and visual acuity

One hundred and four patients (91 probands) with multiple CRB1 variants were included in this cohort and ascertained for phenotyping, after a multidisciplinary team of ophthalmologists and clinical geneticists excluded other possible genotypes. Eighty-seven (84%) were seen at Moorfields Eye Hospital and 17 (16%) at St John of Jerusalem Eye Hospital. Seven patients had a typical phenotype for the disease.
(nummular pigmented deposits, admixed with small yellow-white dots, preserved para-
arteriolar retinal pigment epithelium (PPRPE), abnormally laminated and thickened retina), however a single disease-causing variant in CRB1 was identified. These 7 cases with presumed CRB1-associated disease were excluded from further analysis.

Supplementary Table 1 (available at AJO.com) summarizes the genetics and the clinical phenotype of these patients.

Sixty-two were male (60%) and 42 female (40%). The mean age ± standard deviation (SD) at the first visit was 19.8 ± 16.1 years of age, with a median of 15 years. Fifty-three participants had their first visit as children (below the age of 16). The mean follow-up time of the cohort was 9.6 ± 10 years and the overall age at their latest visit was 29.6 ± 17.2 years of age. The clinical findings from the cohort are summarized in Table 1.

Based on ophthalmic history, imaging, and clinical examination, 54 presented with EOSRD/LCA (51%), 26 individuals were diagnosed with RP (26%), and 24 with MD (23%). Within the EOSRD/LCA sub-cohort, the mean age of onset was 2.4 ± 2.8 years old (median 1), with poor central vision and secondly nystagmus as the most prevalent initial symptoms/signs. Among the RP group, the mean age of onset was 13.2 ± 10.8 years old (median 10) and the most common presenting symptom was nyctalopia, followed by constricted field. Lastly, the patients from the MD sub-cohort had a mean age of onset of 16.8 ± 10.8 years of age (median 15) and primarily complained of decreased acuity.

Baseline BCVA was 1.6 ± 0.8 LogMAR in those with EOSRD/LCA (mean age 16.2 ± 15.3 years), 0.9 ± 0.8 LogMAR in patients with RP (mean age 23.9 ± 18.6 years), and 0.6 ± 0.4 LogMAR in MD patients (23.2 ± 13.7 years). Seven infants with EOSRD/LCA did not have accurate vision recorded (e.g., fix and follow) and 1 patient with RP did not have BCVA detailed in the medical records. The number of patients in each WHO category of visual impairment is displayed in Table 1 and Supplementary Figure 1A (available at AJO.com). Asymmetric BCVA was seen in 20 patients (19%); 9 with EOSRD/LCA, 5 with RP, and 6 with MD. There was a significant association between age and BCVA in all EOSRD/LCA (p <0.0001), RP (p <0.0001), and MD sub-cohorts (p = 0.047).
The mean spherical equivalent in the RP cohort was $+1.75 \pm 1.75$ (n=1 myopic); in the EOSRD/LCA, $+5.75 \pm 3.5$ (n=2 myopic), and $+0.75 \pm 2.5$ among the MD sub-cohort (n=7 myopic, 29%). High hyperopia (spherical equivalent $>5.00$ dioptres) was found in 1 patient with RP, 1 with MD, and in 21 with EOSRD/LCA (39%).

**Clinical findings - anterior segment**

One individual with RP and four with EOSRD/LCA were diagnosed with keratoconus. Fifteen patients with RP had lens opacities (58%), diagnosed at 34 ± 13 years of age; 17 patients among the EOSRD/LCA group (31%) at age 34.5 ± 14; and only two patients in the MD cohort, at ages 54 and 71 respectively. Only four patients among the RP sub-cohort had glaucoma; one neovascular at age 41, two open angle at ages 32 and 35, and one acute angle closure at 29 years of age.

**Clinical findings - posterior segment**

All EOSRD/LCA patients presented with diffuse, dense pigment in the retinal periphery, with both spicules and nummular pigment. They all had macular involvement, with a pigmented ring in the posterior pole in 16 patients, and coloboma-like severe atrophy in four (Figure 2A and B). All patients in the RP sub-cohort presented with peripheral retinal pigmentation and a range of phenotypes at the posterior pole: seven patients had a normal appearing macula, ten had a blunted/opaque macular reflex, and 12 had signs of atrophy and pigmented deposits (Figure 2C and D). Sixteen patients with MD had normal retina outside the arcades, whereas eight had peripheral areas of pigmented bone spicules or nummular lesions, and/or vessel thinning. The macula reflex was blunted in 12 patients, with mottled RPE and signs of atrophy in the remaining 12; the latter often accompanied by pigmented bone spicules and affecting the nasal peripapillary area (Figure 2E and F). Four individuals with MD presented with foveal sparing and therefore good central vision. Macular involvement (functional and/or
structural) was present in 25 RP patients (first documented at mean age of 23.9 ± 18.6 years), and in all EOSRD/LCA (2.4 ± 2.8 years) and MD patients (19 ± 10 years). Thirty-four individuals with EOSRD/LCA (mean age 24.1 ± 16.6 years), 11 with RP (23 ± 18 years), and 3 with MD (37.3 ± 11.1 years) had nummular pigment clumps. White/yellow dots were seen in 16 EOSRD/LCA patients (mean age 24 ± 9 years), in 3 patients with RP (19 ± 7.5 years), and in 2 MD patients, at the same timepoint where nummular pigment was present (25 and 35 years of age). Sixteen patients with EOSRD/LCA (mean age 22 ± 13.5 years) presented with PPRPE; ten with RP (23 ± 12 years), and three with MD (38 ± 10.5 years). PPRPE was more readily identified with autofluorescence (AF) imaging, likely given the RPE-basis of AF imaging.\textsuperscript{10}

Seven EOSRD/LCA patients had retinal telangiectasia (mean age 30 ± 10 years), six of which were associated with exudation, with two resulting in vitreous haemorrhage(s). Retinal telangiectasia was seen in 4 patients with RP, diagnosed at a mean age of 28 ± 8 years, and resulted in a range of complications including exudation, vitreous haemorrhage, and retinal detachment (RD). Optic disc drusen was the most common optic nerve abnormality; affecting 7 patients with LCA, 4 with RP, and 1 with MD. Two patients with EOSRD/LCA had gliosis on the optic disc.

Ocular complications were seen in twelve individuals with EOSRD/LCA (21%); in 4 they were related to telangiectasia, while the remaining eight corresponded to RD, iris nodules, anterior segment synechiae, retinal hamartoma, and corneal hydrops. Ocular complications were also reported in six patients with RP (22%); 4 were associated with retinal telangiectasia, the remaining two were uveitis and asteroid hyalosis. Only one patient with MD was found to have unusual vessel sheathing at age 8, of unknown cause.

**Macular OCT analysis**

Eighty-five patients from our cohort had macular OCT scans (82%); 38 (45%) with EOSRD/LCA, 25 (29%) with RP, and 22 (26%) with MD; with all being included in the qualitative analysis. Quantitative assessment was possible in 15 patients with
EOSRD/LCA, 8 with RP, and 10 with MD, due to image quality and sufficient scans being available. Eleven individuals from the EOSRD/LCA sub-cohort had follow up scans (over 7 ± 3 years), 7 from the RP group (over 7.5 ± 1.52 years), and 10 with MD (7 ± 3 years), which enabled additional longitudinal analysis. Baseline and follow up structural parameters are described on Supplementary Table 2.

The EOSRD/LCA sub-cohort had significantly increased inner ring thickness (IRT, \( p = 0.002 \)), outer ring thickness (ORT, \( p < 0.0001 \)), inner ring volume (IRV, \( p = 0.002 \)), and outer ring volume (ORV, \( p = 0.0004 \)) compared to the normal population (Figure 3). The RP group had significantly decreased central macular thickness (CMT, \( p = 0.0008 \)) and central macular volume (CMV, \( p = 0.0037 \)), however the ORV was increased (\( p = 0.0004 \)). Lastly, patients with MD showed significant thinning in all CMT (0.003), IRT (0.004), ORT (0.01), CMV (0.0017), and IRV (0.01; Figure 3A). No association was found between thickness or volume metrics and age in EOSRD/LCA (\( p \) between 0.08 – 0.95) and MD subgroups (\( p \) between 0.4 and 0.8). However, a significant association was found in patients with RP between age and ORT (negative slope, \( p = 0.003 \)) and IRV (positive slope, \( p = 0.0003 \), Figure 3B). Longitudinal analysis demonstrated no statistically significant differences between baseline and follow up parameters in all subcohorts. Sixty percent of patients with EOSRD/LCA had decreased CMT over follow up (mean -3 \( \mu \)m/year), 54% had decreased IRT (-1.5 \( \mu \)m/year), and 50% had decreased ORT (-0.6 \( \mu \)m/year). Within the RP sub-cohort, 67% had decreased CMT (-1.9 \( \mu \)m/year), 49% had decreased IRT (with an overall increase however of 0.3 \( \mu \)m/year), and 43% had decreased ORT (increase of 1.5 \( \mu \)m/year). Lastly, 59% patients with MD had decreased CMT over follow up (-0.5 \( \mu \)m/year), 70% had decreased IRT (-1.1 \( \mu \)m/year), and 49% had decreased ORT (-1.8 \( \mu \)m/year).

Macular cystic spaces were present in 29% of our cohort as a whole, and in 52% and 42% of the RP and MD subgroups, respectively. The majority had the INL affected only (24 eyes), followed by both INL and ONL involved (19 eyes), and then ONL only (13 eyes). There was a significant difference in the age of patients with and without cysts, both analysing the cohort as a whole (0.007) and excluding the EOSRD/LCA group (\( p = 0.002 \)), with a mean age of 22.3 ± 15.4 for those having cysts and 34.6 ± 18.2 for those
The cysts resolved during follow up in 24% of eyes, without any treatment. The presence of outer layers at the fovea and perifovea was also recorded (supplementary results).

In terms of qualitative assessment, overall, nine patients had both eyes in OCT group 1 (11%, 28.9 ± 16.4 years old), 52 in group 2 (61%, 25.2 ± 15.4 years old), and 17 in group 3 (20%, 43.2 ± 14.8 years of age, 15 of which were in the EOSRD/LCA cohort; Figure 1 and Supplementary Figure 1B). Two patients had one eye in group 1 and the other in group 2; and five patients had one eye in group 2 and the other in group 3.

Considering the cohort as a whole, there was a significant age difference between groups 1 and 3 (p 0.03) and 2 and 3 (0.0003), while no age difference was recorded between groups 1 and 2 (p 0.49). Visual acuity was also significantly different between groups; LogMAR 0.4 ± 0.32 group 1, 1.25 ± 0.97 group 2, and 3.27 ± 1.38 group 3; with progressive VA decline as the OCT layers became more disorganised (p <0.0001).

Within the EOSRD/LCA group, 20 patients had both eyes in OCT group 2 (51%, 19.6 ± 11.1 years old), 16 in OCT group 3 (42%, 38.4 ± 16.9 years old), and two each eye in groups 2 and 3 (ages 15 and 27). The RP sub-cohort had three patients in group 1 (12%, 34 ± 14 years old), 17 in group 2 (68%, 22.8 ± 17.9 years old), two in group 3 (44 and 53 years old), one with one eye in group 1 and the other in group 2 (37 years old), and two with one eye in group 2 and the other in group 3 (38 and 73 years old). Lastly, the MD group had five patients in group 1 (23%, 16.3 ± 17 years old), 15 in group 2 (68%, 31.7 ± 15.8 years old), one with one eye in group 1 and the other in group 2 (15 years old), and one with one eye in group 2 and the other in group 3 (23 years old).

Findings summarised in Table 1. Focusing on the EOSRD/LCA sub-cohort, the age difference between groups 2 and 3 was significant (p 0.0005), while no age difference was found between groups 1 and 2 of the RP (p 0.28) and MD sub-cohorts (p 0.07).

In addition, given image quality variation, potential challenges to robustly distinguish between OCT groups 2 and 3, and the desire to compare definitively normal with abnormal architecture, we also compared group 1 (normal lamination) vs groups 2 and 3 combined (abnormal lamination). The latter (group 2+3) had a mean age of 26.9 ± 16.8 years old and there was no significant age difference when compared to the
normal lamination group (p 0.81). The visual acuity of the abnormally laminated group was LogMAR 1.75 ± 1.39, which was significantly different to group 1 (p <0.0001).

In terms of transition between groups over time: 4 eyes (3 patients) with EOSRD/LCA changed from group 2 to 3; while five (3 patients) with RP, and five (3 patients) with MD went from group 1 to group 2 during follow up (further details in Supplementary Results, available at AJO.com).

**Electrophysiology**

Twenty-nine patients (26%) underwent ff-ERG and PERG. There was a high degree of inter-ocular ERG symmetry based on amplitudes of the DA 0.01, DA 3 and DA 10 ERG a- and b-waves, LA 30Hz ERG and LA 3 (single flash) ERG b-waves (slope = 1.00; r²= 0.98), and on the peak times of the DA 10 ERG a- and b-waves and LA 30Hz ERGs (slope = 0.98; r²=0.96). PERG N95 data was available for analysis in 12 of 14 patients with a detectable P50 component. The N95:P50 ratio had a mean value of 1.7 (range 1.4 to 3.0), which is similar to the reference group for the lab (mean = 1.5); i.e. there was no evidence of N95 reduction disproportionate to P50 in this small group.

*Figure 4* summarises the electrophysiological findings and Supplementary Figure 2 shows representative recordings. Eight patients with a clinical diagnosis of EOSRD/LCA aged between 9 and 27 years had undetectable ERGs under all stimulus conditions. The ERGs in others (8 patients with MD and 6 with RP) revealed a similar degree of rod and cone system involvement, although there was evidence of slightly greater rod than cone system dysfunction in 5 cases, revealed by proportionately greatest reduction in the rod-system selective DA 0.01 ERG (Figure 5a; patients 8, 13, 14, 16 and 19). Seven patients had MD both clinically and electrophysiologically, including 6 with normal ERGs and one with mild reduction in the LA 30z ERG (*Figure 4A*, patient 28), but likely due to eye closure noted during testing. Further details are described on Supplementary Results and Supplementary Figures 2 and 3.

**Longitudinal analysis**
Follow up BCVA was obtained in 41 (76%) individuals with EOSRD/LCA, 20 (77%) with RP, and 20 (83%) with MD. Mean follow up time was 9.9 ± 12.1 years for patients with EOSRD/LCA, 9.9 ± 9.8 for the RP sub-cohort, and 7.6 ± 5.7 for MD patients. Follow up BCVA was 1.9 ± 0.7 LogMAR in patients with EOSRD/LCA (mean age 27 ± 17 years), 1.3 ± 1 LogMAR in patients with RP (38 ± 13 years old), and 0.8 ± 0.5 LogMAR in MD patients (30 ± 16 years). There was a significant difference between baseline and follow up BCVA in the EOSRD/LCA (p <0.0001), RP (p <0.0001), and MD (p =0.016) sub-cohorts.

The rate of BCVA decline was 0.06 LogMAR (3 letters)/year in patients with EOSRD/LCA, 0.07 LogMAR (3.5 letters)/year in patients with RP, and 0.04 LogMAR (2 letters)/year in patients with MD. Fifteen patients (35%) with EOSRD/LCA dropped 15 ETDRS letters or more over follow up in at least one eye, 10 (45%) with RP, and 7 (35%) with MD. Fourteen patients (33%) with EOSRD/LCA progressed to more advanced WHO categories of visual impairment, 7 (32%) with RP, and 6 (30%) with MD. Overall, 20 patients (24%) became severely sight impaired and/or blind over follow up; 12 with EOSRD/LCA, 5 with RP, and 3 with MD.

Macular OCT longitudinal evaluation demonstrated that EOSRD/LCA patients may experience slowly decreasing retinal thickness -despite their baseline thickening-, progressive loss of lamination, and degeneration of both central and pericentral outer retinal layers over follow up. CRB1-RP patients in our sample experienced loss of thickness at the fovea and parafovea, while increasing in the perifovea; although the small sample size available for longitudinal assessment should be borne in mind. MD patients had progressive thinning of the posterior pole, associated with poor retinal lamination, and discontinuous/narrower outer layers.

**Genetics**

Seventy-eight different CRB1 variants were found in our cohort, 46 of which were missense (59%), 11 frameshift (14%), 10 nonsense (13%), 5 canonical splice site alteration (6%), 2 missense with significant splice site alteration (3%), 2 intronic variants
(3%), 1 in-frame deletion (1%), and 1 deletion of exon 12 (1%) (Supplementary Table 3). Schematic presentations of these detected variants and the status of evolutionary conservation are presented (Figure 5A, Supplementary Figures 4 and 5). Twenty-eight variants were classified as pathogenic (36%), 43 (56%) were likely pathogenic, and 7 (8%) were variants of uncertain significance. Twenty-three variants (29%) were novel, whereas 55 were previously reported.

c.2843G>A, p.(Cys948Tyr) was the most common variant, seen in 15 individuals (14 probands), 14 of which had EOSRD/LCA. This was followed by c.498_506delAATTGATGG, p.(Ile167_Gly169del), found in nine probands with MD. Another frequent variant among the MD group was p.Pro836Thr, present in seven black African individuals (4 families) with MD.

Null variants (including nonsense, frameshift, splicing, and exon deletion) were significantly more associated with both EOSRD/LCA (p 0.001) and MD (p 0.003) phenotypes, rather than with RP (Figure 5B). Double null genotypes were only seen in EOSRD/LCA (12 patients). The four variants p.Cys896Ter, p.Thr745Met, p.Ser478ProfsTer24, and p.Cys195Phe; and the three loci p.Gly770, p.Gln1219, and p.Pro1381 harboured changes causing both EOSRD/LCA and MD (Figure 4A). In the MD cases however, the aforementioned variants and loci were combined either with p.(Ile167_Gly169del) - previously characterised as a hypomorphic allele; or with a likely pathogenic variant. Eight individuals with MD (32%, 7 probands) did not harbour the in-frame or the p.Pro836Thr variants. p.Asn880Ser was the only repeated variant among the latter, seen in two unrelated families.

Intrafamilial variability was present in two families. In one family, a sibling had mild, asymptomatic MD and the other had RP with decreased vision since her 20s (homozygous p.Ser638Leu). In the other family, three different CRB1 variants were found to be segregating; a great uncle inherited p.Cys195Phe and p.(Ser478ProfsTer24) and developed EOSRD/LCA, and one nephew harboured p.(Ser478ProfsTer24) and the in-frame p.(Ile167_Gly169del) and presented with MD.
DISCUSSION

This study characterises the largest cohort of patients with molecularly confirmed CRB1-retinopathy to date, with 23 novel CRB1 variants identified. Deep phenotyping data is presented, including detailed analyses of demographic features, fundus imaging, morphological characteristics, and quantitative data of electrophysiological assessment, with the aim of establishing clear phenotypes, potential outcome measures, genotype-phenotype correlations, and laying the foundations for future studies that aim to develop therapies and treatment strategies.

Low vision was the most common WHO category for individuals with EOSRD/LCA in the first two decades of life, followed by blindness from 20 years on. In the RP group there was severe visual impairment from 40 years of age, while in the MD group it was only seen in a minority of cases from 28 years old. This is similar to the observations from Talib et al., who reported blindness among CRB1-RP patients from the fifth decade of life. Patients with MD appeared to have an overall good prognosis, with relatively preserved visual function until adult age in at least one eye. The findings highlight the window of opportunity for potential treatments such as gene therapy. Individuals with EOSRD/LCA may need relatively early intervention in childhood and adolescence due to early macular structural and functional involvement, whereas patients with RP or MD would potentially benefit up until their late 30s. The latter is in agreement with Talib and Mathijssen et al., while Nguyen et al. recommended treatment before the third decade of life, having seen in their cohort a faster decline around that period.

Furthermore, in our cohort, VA deterioration was significantly associated with age in all EOSRD/LCA, RP, and MD sub-cohorts. This association was also found in CRB1-RP by Mathijssen et al., but not by Talib et al., despite their cohort being mostly RP phenotype. Employing their statistical approach of age vs BCVA of the best seeing eye (instead of age vs BCVA of both eyes as we have herein), we still obtained a significant association (p=0.001). Regarding BCVA asymmetry, we report a smaller proportion than previous cohorts (19% in this report vs 33% and 31% in other groups). Progressive visual decline was also reported during long follow up studies, however this was not picked up in the 2-year follow up study by Nguyen et al.
The rate of progression of our patients was greater than the one reported by Talib et al., possibly reflecting an overall earlier onset and more severe disease stage in our patients.

The median age of onset in the RP subgroup was 10 years, which was older than the Dutch and Belgian cohorts (4 and 5 years each). Therefore, 45% of our RP group had a disease onset after the first decade of life (versus 15% and 20% in the aforementioned cohorts). We appear to have captured a subset of individuals with mild, adult-onset disease, previously poorly characterised. Seven of these patients (14 - 53 years of age) had preserved foveal architecture and good central vision at their latest visit.

Hyperopia was the prevalent refractive error in all CRB1 phenotypes, with 37% of the EOSRD/LCA patients having high hyperopia (>+5.00 D). Wang et al. found over 80% of their CRB1-LCA cohort to be highly far-sighted and Mathijssen et al. also found consistent hyperopia in their CRB1-RP cohort. This was not however a pathognomonic feature in CRB1-MD (28% of our patients were myopic and 16% were emmetropic), corroborating a previous study.

The most prevalent posterior segment feature within our cohort was maculopathy, which was present in 97% of our cohort, with variable severity, and consistent with smaller studies. Other common features included nummular pigment, PPRPE, white/yellow dots, and retinal telangiectasia. We did not find significant differences between the ages at which white/yellow dots and nummular pigment were found, and sometimes both appeared simultaneously (Figure 2A), therefore it is unclear if these represent different stages of the same lesion. Of note, pigmented para-venous chorioretinal atrophy, previously associated with CRB1 on one occasion, was not seen in any of our genetically confirmed patients.

Macular OCT revealed a range of features; 51% of the eyes from our cohort had ill-defined lamination (group 2; Figure 1), and 18% had disorganised retinal layers (group 3). Previous studies have reported similar values, although these may depend on the cohort characteristics e.g., EOSRD/LCA patients have a more severe phenotype. Our results are comparable with some previous investigations e.g., where 38 to 52% eyes
were categorised as group 3.\textsuperscript{37,45} However, we consider that the classification of OCTs into groups 2 and 3 can be somewhat subjective and it is also dependant on the quality of the scan, intrinsically difficult in patients with EOSRD/LCA and nystagmus. To mitigate this potential limitation and thereby minimize possible bias, we added the comparison of normal and abnormal lamination groups. Patients in group 3 were older and our longitudinal data (available in 7\% of eyes) showed progressive loss of defined lamination on qualitative review; therefore, the poorly laminated and disorganised OCT in \textit{CRB1} may also involve a degenerative process, and not only a congenital one. This is consistent with previous suggestions; although far larger studies with greater serial data is required.\textsuperscript{46} However, it is of note, that an age difference was not present when comparing the groups of normal versus abnormal OCT lamination. Nevertheless, given its association with poor visual function, OCT layer organisation should be considered as a criterion to take into account for future treatment trials.\textsuperscript{36,37} Degeneration of central and pericentral outer layers during follow up was also reported by other groups in longitudinal analysis.\textsuperscript{10}

Quantitative analyses of OCT have revealed a strong association between CRB1 and a thickened retina, reported in animal models\textsuperscript{49} as well as in up to 82\% of the individuals from a previous cohort.\textsuperscript{46} The perifovea (outer ETDRS ring) has been the most commonly described area with increased thickness, with limited evidence regarding the central and inner ring areas.\textsuperscript{10,14,20,46} We observed that our EOSRD/LCA and RP groups had significantly increased outer ring thickness, when compared to the reference population. We also noticed increased inner ring values in our EOSRD/LCA cohort exclusively, previously unreported; while patients with \textit{CRB1}-MD seemed to be exempt from the retinal thickening characteristic. Some groups have also reported age-dependant posterior pole thinning.\textsuperscript{10,20} Although we found decreased macular thickness longitudinally in our EOSRD/LCA and MD sub-cohorts, this did not reach statistical significance. We found a significant decrease in outer ring thickness with age, concomitantly with increased inner ring volume in the RP subgroup. It is possible that the thinning follows a centripetal pattern and the parafovea remains preserved until later stages; yet, a larger number of patients is needed to arrive at a more definitive conclusion. The inner retina is generally where the increased thickness is most
noticeable, which may be due to excess ganglion cells. Our hypothesis is that increased retinal thickness is due to CRB1-related poor adhesion between cells, which leads to poor lamination and abnormal anatomy, consequently occupying volume in a less efficient, broader way. 

Macular cystic spaces were present in a lower percentage in our cohort when compared to other groups, and this may be related to the higher proportion of EOSRD/LCA patients, who are rarely affected by macular fluid accumulation. We were able to associate younger age with the presence of macular cystic spaces, a relationship that was previously hypothesized. Cysts resolved in nearly a quarter of the eyes during follow up without any treatment, which is noteworthy given their reported poor responsiveness to treatment. We can conclude that cystoid maculopathy is a prevalent feature in CRB1-RP and -MD, affecting mostly younger individuals, and it can be self-limiting. This has caused CRB1 to be included within the differential diagnoses considered in cystic maculopathies, such as those secondary to variants in RS1 and NR2E3. Although macular fluid accumulation is also seen in other RP genotypes, the pathophysiology may be easier to explain in CRB1-retinopathy. CRB1 transcript CRB1-A has been located within Müller glia cells in mouse and human retina, its loss manifesting as an irregular number and size of their apical villi. Müller cell loss of integrity may lead to abnormal fluid transport, and has been associated with macular oedema; hence this may be the cause of the cystic spaces found in some CRB1 phenotypes.

The variant c.2843G>A has been reported the most commonly occurring in individuals with CRB1-related disease, having a carrier frequency of approximately 2:10,000. c.2843G>A, p.(Cys948Tyr) was also the most prevalent in our cohort, as well as in other European groups. However, in a Dutch cohort, p.(Met1041Thr) was the most prevalent, and in a Chinese cohort p.(Gly122Ter) was most common, and the c.2843G>A allele was not identified. The in-frame deletion p.(Ile167_Gly169del) was reported to be the most prevalent disease-causing CRB1 variant in non-Asians, particularly frequent in Spanish patients. It was previously described as a hypomorphic
allele, which led to limited retinal dysfunction. However, two Spanish reports related it to early onset RP and LCA, without further phenotypic details. We found p.(Ile167_Gly169del) in nine patients with MD. Eighty percent of these patients had cystic macular changes, and 90% did not have visual impairment as per WHO classification. Therefore, this variant may be associated with a good clinical prognosis, with possibly limited retinal involvement (yet not necessarily macula restricted).

Another frequent variant among the MD group was p.(Pro836Thr), present in seven black African individuals (4 families). This variant was first described by den Hollander et al., seen in a patient with RP and PPRPE, without a known second variant, and with no details regarding race. It was also reported in two identical twin sisters with cystic maculopathy. It has a frequency of 0.00277 and 0.0028 in the black African population or descendants (an allele frequency of exome and genome: gnomAD v2.1.1), and it currently has conflicting evidence regarding its pathogenicity (ClinVar ID: 372352). The new cases associating it with disease, both in compound heterozygous and homozygous state, contribute towards classifying it as disease-causing.

Lastly, we were able to test the association of null variants and different phenotypes. We not only found them significantly associated to EOSRD/LCA, as previously suggested by other groups, but also to MD, representing 36 and 26% of the variants seen in each group. In contrast however, null variants in the MD sub-cohort were combined with likely hypomorphic / milder pathogenic variants, thereby decreasing the overall functional impact of these genotypes. Also, the relatively small number of MD patients compared to EOSRD/LCA patients in our cohort needs to be borne in mind.

The large size of CRB1, occupying nearly all the AAV packing capacity, has made the development of a therapy for these retinopathies challenging. However, novel approaches through AAV vectors with small promoters have allowed the expression of CRB1 protein in Müller glial cells in vitro. Another option tested in vivo was the supplementation of CRB2, a smaller yet highly similar protein in structure, that improved the photoreceptors' morphology and function after delivery. These creative approaches, as well as the innovative human induced pluripotent stem cells (iPSCs) disease models, will facilitate progress to developing treatment options.
the detailed characterization of CRB1 macular OCT features will likely be helpful towards its inclusion among the inherited retinal dystrophy genotypes that are currently used to train and validate deep learning systems for automated diagnosis and classification.\textsuperscript{68}

Inherent limitations of our study are its retrospective design, the differences in follow up time between patients, not all data being available for every individual, and different types of accessible data, acquired with diverse methods and protocols. The above-mentioned limitations were ameliorated by the size of our genetically-confirmed cohort, the wide age range of our patients, the international representation, and their detailed medical records.

In conclusion, this multinational investigation characterises the largest cohort of patients with molecularly confirmed CRB1-retinopathy to date, identifies novel CRB1 variants, and delineates genotype-phenotype correlations. Clinical and functional phenotypes are detailed, establishing the range and combinations of features that will aid diagnosis and inform patient management. Potential endpoints for future natural history and interventional clinical trials are highlighted, laying the foundations for future studies that aim to develop therapies and treatment strategies.
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FIGURE LEGENDS

Figure 1. Macular OCT findings and qualitative classification. A) Group 1, characterized by normal lamination; to the left it corresponds to a 16-year-old patient with MD and macular cystic spaces, and to the right to a 33-year-old patient from the RP subgroup. B) Group 2, where the retinal layers are still discernible but appear ill-defined; on the right the image corresponds to a 50-year-old individual with MD, and on the left an 11-year-old girl with LCA and macular cystic spaces. C) Group 3, defined by a disorganised retina with coalescent layers (particularly within the inner retina) and increased reflectivity of the nuclear layers; the images correspond to two patients with EOSRD/LCA, at age 27 years to the left and 29 years old to the right.

Figure 2. Ultra-widefield color and autofluorescence fundus images from individuals with CRB1-retinopathy. A) Thirty-year-old patient with EOSRD/LCA, his parents and paediatrician noticed he had nystagmus as an infant and he had poor vision his entire life. Deep pigmented nummular lesions, as well as yellow/white dots mainly temporal to the macula, are seen. Autofluorescence (AF) is decreased and an optic disc drusen is visible. B) Sixty-one-year-old patient with LCA. More coalescent pigment deposits and profound macular atrophy are seen. In the AF image preserved para-arteriolar retinal pigment epithelium (PPRPE) is more readily seen. C) Twenty-eight-year-old with RP and retinal telangiectasia which resulted in exudation and vitreous haemorrhages. D) Eighteen-year-old with RP, with few pigment deposits, macular involvement and PPRPE. E) Seventy-year-old patient with MD, he reported decreased central vision since age 15 years and normal peripheral field. F) Forty-two-year-old with MD, he had failed a vision screening test at age 24 years and his vision had been slowly decreasing since. Of note is the characteristic pattern of hypoautofluorescence that involves the macula and all optic disc borders.

Figure 3. Graphic representation of statistical analysis undertaken to analyse various structural OCT parameters. A) Bar graphs comparing structural measurements in each group to a control group. Significant differences in all groups are seen (marked with *), with the EOSRD/LCA group showing significantly increased inner ring thickness (IRT), outer ring thickness (ORT), inner ring volume (IRV), and outer ring volume (ORV); the
RP group with decreased central macular thickness (CMT) and central macular volume (CMV), and increased ORV; and the patients with MD with thinned CMT, IRT, ORT, CMV, and IRV. B) Linear regression representation of the association between age and structural parameters in the RP sub-cohort, where we see a significant, positive slope regarding IRV, and a negative one when it comes to ORT.

**Figure 4.** Quantification of the full-field and pattern electroretinogram (ERG; PERG) findings. Full field ERG findings summarised in 29 subjects tested according the ISCEV standard methods; a) The amplitudes of the DA 10 ERG a-wave, LA 30 Hz ERG and LA 3 ERG b-wave are plotted against the primary axis as a percentage of the age-matched lower limit of the (“normal”) reference range, with values arranged in ascending order of DA10 ERG a-wave amplitude for clarity. The LA 30 Hz peak times are plotted as a difference from the age-matched upper limit of normal timing against the secondary axis. b) Pattern ERG P50 amplitudes plotted as a percentage of the lower limit of normal amplitude for the same order of subjects as in a); the absence of a column indicates undetectable responses. c) The age of the patients at the time of testing, arranged in same order as in a) and b). Note the 7 subjects to the right of the vertical broken line in a) had normal ERGs but abnormal pattern ERG P50 components (b), consistent with a diagnosis of macular dystrophy. The LA 30Hz ERG in subject 29 was subnormal, but in presence of significant eye closure.

**Figure 5.** A) Graphic representation of the **CRB1** gene and protein, with details on functional domains. Each variant within our cohort is detailed above the protein, with its corresponding location, and with different colors according to the associated phenotype. B) Pie graph representation of the type of variants seen in each phenotypic group. The larger prevalence of null changes can be easily visualized in the MD and EOSRD/LCA groups, compared to the RP group, and also the high prevalence of the p.Ile167_Gly169del in-frame deletion, found almost exclusively in the MD group.
Table 1: Clinical characteristics of CRB1 disease.

<table>
<thead>
<tr>
<th></th>
<th>EOSRD/LCA (n=54, 52%)</th>
<th>RP (n= 26, 25%)</th>
<th>MD (n= 24, 23%)</th>
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<tbody>
<tr>
<td>Age at baseline, mean ± SD (years)</td>
<td>16.2 ± 15.3</td>
<td>23.9 ± 18.6</td>
<td>23.2 ± 13.7</td>
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<tr>
<td>Gender, n (%)</td>
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<tr>
<td>Male</td>
<td>32 (56)</td>
<td>19 (66)</td>
<td>17 (68)</td>
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<tr>
<td>Female</td>
<td>25 (44)</td>
<td>10 (34)</td>
<td>8 (32)</td>
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<td>Age of onset, mean ± SD (years)</td>
<td></td>
<td></td>
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<tr>
<td>Infancy (birth – 2 years old)</td>
<td>37</td>
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<td>0</td>
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<tr>
<td>Childhood (3 – 11 years old)</td>
<td>20</td>
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<td>10</td>
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<tr>
<td>Adolescence (12 – 16 years old)</td>
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<td>5</td>
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<td>Adulthood (over 16 years old)</td>
<td>0</td>
<td>10</td>
<td>9</td>
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<tr>
<td>Baseline best corrected visual acuity, mean ± SD (LogMAR)</td>
<td>1.6 ± 0.8</td>
<td>0.9 ± 0.8</td>
<td>0.6 ± 0.4</td>
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<tr>
<td>Final best corrected visual acuity, mean ± SD (LogMAR)</td>
<td>1.9 ± 0.7</td>
<td>1.3 ± 1</td>
<td>0.8 ± 0.5</td>
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<td>Baseline WHO visual impairment category, n= (%)*</td>
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<tr>
<td>No or mild impairment</td>
<td>4 (7)</td>
<td>10 (38)</td>
<td>14 (58)</td>
</tr>
<tr>
<td>Moderate impairment</td>
<td>15 (28)</td>
<td>10 (38)</td>
<td>10 (42)</td>
</tr>
<tr>
<td>Severe impairment</td>
<td>8 (15)</td>
<td>0</td>
<td>0</td>
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<tr>
<td>Blindness</td>
<td>20 (37)</td>
<td>5 (21)</td>
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<td>Spherical equivalent, mean ± SD</td>
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<tr>
<td>High hyperopia</td>
<td>21 (39)</td>
<td>1 (3)</td>
<td>1 (4)</td>
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<tr>
<td>Myopia</td>
<td>2 (4)</td>
<td>1 (3)</td>
<td>7 (29)</td>
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<td>Lens opacity, n (%)</td>
<td>17 (31)</td>
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<td>Keratoconus, n (%)</td>
<td>4 (7)</td>
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<td>Nummular pigment, n (%)</td>
<td>34 (63)</td>
<td>11 (42)</td>
<td>3 (12)</td>
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<td>Macular involvement, n (%)</td>
<td>54 (100)</td>
<td>25 (96)</td>
<td>24 (100)</td>
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<td>Baseline OCT categories, eyes n (%)</td>
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<td>Normal lamination and organisation</td>
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<td>Abnormal lamination</td>
<td>42 (54)</td>
<td>37 (75)</td>
<td>32 (70)</td>
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<td>Disorganisation</td>
<td>34 (46)</td>
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<td>1 (2)</td>
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<td>Yellow/white dots, n (%)</td>
<td>16 (30)</td>
<td>3 (11)</td>
<td>2 (8)</td>
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<td>PPRPE, n (%)</td>
<td>16 (30)</td>
<td>10 (38)</td>
<td>3 (12)</td>
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<tr>
<td>Retinal telangiectasia, n (%)</td>
<td>7 (13)</td>
<td>4 (15)</td>
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EOSRD/LCA: Early Onset Severe Retinal Dystrophy/ Leber Congenital Amaurosis; RP: Retinitis Pigmentosa; MD: Macular Dystrophy; SD: Standard Deviation; OCT: Optical Coherence Tomography; PPRPE: Preserved Paraarteriolar Retinal Pigment Epithelium.
*: the patients that do not appear in this classification correspond to 7 babies with LCA with non ETDRS measured vision (e.g., fix and follow) and 1 patient with RP with no details of acuity.

This is multicentre international retrospective cohort study.

Clinical notes, ophthalmic images, and genetic testing results of 104 patients with disease-causing CRB1 variants were reviewed.

EOSRD/LCA is the most common phenotype, often associated with null variants.

Individuals with CRB1 variants can present with adult-onset RP.

OCT lamination may have a degenerative component.

CRB1-retinopathy is often symmetric and has a reasonable window for intervention.