Clinical exome sequencing for inherited retinal degenerations at a tertiary care center

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Inherited retinal degenerations are clinically and genetically heterogeneous diseases characterized by progressive deterioration of vision. This study aimed at assessing the diagnostic yield of exome sequencing (ES) for an unselected cohort of individuals with hereditary retinal disorders. It is a retrospective study of 357 unrelated affected individuals, diagnosed with retinal disorders who underwent clinical ES. Variants from ES were filtered, prioritized, and classified using the ACMG recommendations. Clinical diagnosis of the individuals included rod-cone dystrophy (60%), macular dystrophy (20%), cone-rod dystrophy (9%), cone dystrophy (4%) and other phenotypes (7%). Majority of the cases (74%) were singletons and 6% were trios. A confirmed molecular diagnosis was obtained in 24% of cases. In 6% of cases, two pathogenic variants were identified with phase unknown, bringing the potential molecular diagnostic rate to ~ 30%. Including the variants of uncertain significance (VUS), potentially significant findings were reported in 57% of cases. Among cases with a confirmed molecular diagnosis, variants in EYS, ABCA4, USH2A, KIZ, CERKL, DHDDS, PROM1, NR2E3, CNGB1, ABCC6, PRPH2, RH0, PRPF31, PRPF8, SRNRNP200, RP1, CHM, RPGR were identified in more than one affected individual. Our results support the utility of clinical ES in the diagnosis of genetically heterogeneous retinal disorders.

Retinal diseases are a class of clinically and genetically heterogeneous disorders. To date, more than 300 genes have been described to cause non-syndromic or syndromic retinal degeneration, and the number is increasing (RetNet: https://sph.uth.edu/retnet/; Accessed 04/30/2022). A confirmed molecular diagnosis is important for clinical management, prognostic assessment and more importantly, treatment prospects such as gene therapy approaches are becoming increasingly available for Inherited Retinal Degenerations (IRDs) including the recently FDA-approved voretigene neparvovec-rzyl (Luxturna) for RPE65 gene-specific Leber Congenital Amaurosis (LCA)1, and clinical trials are ongoing for gene therapies of several other IRD genes. It is likely that within the next few years we will see a rapid expansion of gene specific therapies and treatments for retinal disease3. Given this, the urgency to identify a molecular diagnosis in patients with retinal dystrophy has become critical for personalized medicine, patient care, and the prevention of vision loss in the affected individuals.

Many of the earlier methodologies used for detection of disease causing genetic variation in IRDs such as Sanger sequencing, polymerase chain reaction (PCR) or array-based methods, relied on targeted sequencing of a few genes (gene panels) or variants which were known to be causative for disease. These methodologies were targeted for specific gene/variation with low throughput and yields for molecular diagnosis. With the advent of Next-Generation Sequencing (NGS) based methodologies, analysis of millions of variants in the entire human

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genome became possible in a high-throughput way. This revolutionized the detection of disease associated variation and has led to the rapid expansion of our understanding of the molecular causes of human disease including IRDs, as well as our capability to identify the specific molecular cause of disease.

Given the incredibly heterogeneous nature of retinal disease, NGS based methodologies such as targeted panels or exome sequencing (ES) are ideal for identification of underlying molecular variants associated with this disease phenotype. Targeted capture or virtual panels from ES are able to identify rare molecular variants within a specific subset of genes known to be associated with retinal disease.

Currently for IRD, both targeted panels and ES tests are routinely utilized by laboratories worldwide, and each has benefits and limitations. Targeted panels often have overall better coverage but lack the ability to detect variation in genes not included on the specific panel. The benefit of ES is that almost all the coding regions of the genome can be simultaneously analyzed. However, ES is limited to the coding exons and few bases of the splice junction of a wider variety of genes, but is not able to detect deep intronic variation which can often be included on specific targeted panels. Further, the capture might not be optimal for several regions, including genes of interest. Despite their benefits and limitations, the overall outcomes seem to be similar for both targeted panel and ES, with potentially significant molecular findings typically identified in > 50% of cases in large cohort studies (Supplemental Table 1). In these studies, targeted panel based testing was utilized for a significant subset or the entirety of the cohort, leading to a slight bias for that methodology over ES. Additionally, in many of the published studies, the description of the criteria used to determine a positive molecular diagnosis were lacking, with only few studies using a standardized classification metric for variants or specific evidence to classify these variants as pathogenic or likely pathogenic for disease, or confirm that the identified variant is just a rare benign variant identified in their test.

Here, we add to this body of knowledge with the results of ES performed in a clinical laboratory for 357 consecutive cases with retinal disorders. These individuals were ascertained at the ophthalmology clinic at Columbia University Irving Medical Center (CUIMC) and clinical ES test was performed in the Laboratory of Personalized Genomic Medicine at CUIMC. Variants identified in these cases were curated and interpreted based on the Variant Interpretation Guidelines described by the American College of Medical Genetics (ACMG).

Results

Demographics. In this study, 357 consecutive unrelated probands with retinal disorders. These individuals were ascertained at the ophthalmology clinic at Columbia University Irving Medical Center (CUIMC) and clinical ES test was performed in the Laboratory of Personalized Genomic Medicine at CUIMC. Variants identified in these cases were curated and interpreted based on the Variant Interpretation Guidelines described by the American College of Medical Genetics (ACMG).
A likely molecular diagnosis was made in 22 cases, all of which had two or more pathogenic or likely pathogenic variants identified in a gene associated with autosomal recessive inheritance. This included multiple cases with two or more variants of unknown phase in ABCA4 (5 cases), USH2A (6 cases), EYS (3 cases), single cases with variants in CNGA3, CNGB1, COL18A1, COQ2, NPHP4, PCARE, PDE6B, and RPE65. These were largely proband only cases (19/22; 86.4%), two duo cases in which one parent was unavailable for testing, and a single case with proband and one child with unknown clinical phenotype submitted for testing.
### Table 2. Most frequently identified genes in the dataset (by number of families).

| Gene                      | No. of cases | Inheritance | Phenotype association                                      |
|---------------------------|--------------|-------------|------------------------------------------------------------|
| RPGR                      | 9            | X-linked    | Cone-Rod Dystrophy, Macular Degeneration, Retinitis Pigmentosa |
| PRPH2                     | 6            | AR, AR      | Leber Congenital Amaurosis, Macular Dystrophy, Retinitis Pigmentosa, Choroidal Dystrophy |
| RHO                       | 5            | AR, AR      | Night Blindness, Retinitis Pigmentosa, Retinitis Punctata Albescens |
| ARCA4                     | 4            | AR          | Cone-Rod Dystrophy, Fundus Flavimaculatus, Retinal Dystrophy, Retinitis Pigmentosa, Stargardt |
| EYS                       | 4            | AR          | Retinitis Pigmentosa                                       |
| KIZ                       | 3            | AR          | Retinitis Pigmentosa                                       |
| PRPF8                     | 3            | AR          | Retinitis Pigmentosa                                       |
| USH2A                     | 3            | AR          | Retinitis Pigmentosa, Usher Syndrome                      |

### Table 3. Genes in which variants were identified by phenotype. P/LP Pathogenic/Likely Pathogenic.

| Phenotype                  | No. of cases | No. of cases with at least 1 P/LP | Genes identified with P/LP variants                                                                 |
|----------------------------|--------------|-----------------------------------|-----------------------------------------------------------------------------------------------------|
| Rod-cone                   | 212          | 94                                | ABC4, AHI1, BEST1, BBS1, CEP90, CNGA1, CNGA3, COQ2, EYS, FAM161A, IFT140, IMPG2, KIZ, KLHL7, MYO7A, NR2E3, PCARE, PCDH15, PDE6B, PRPF8, PRPF31, RHO, RP1, RPGGR, SNRNP200, TOPORS, USH2A |
| Macular dystrophy          | 72           | 28                                | ABC4, CNGA3, CRX, EFEMP1, PDE6C, PROM1, PRPF8, PRPH2, RDH12, RP1L1, RPGGR, RS1                      |
| Cone-rod                   | 32           | 12                                | ABC4, CDR1, GUCY2D, NPHP4, PCARE, PROM1, PRPH2, RPGGR, TULP1                                       |
| Cone dystrophy             | 13           | 4                                 | CACNA2D4, GUC1A1A, MAK, OPN1MW                                                                  |
| Other                      | 29           | 9                                 | ABC6C, CACNA1E, CHM, EYS, GPR143, LCA5, SLC38A8                                                   |

### Potentially clinically significant molecular findings.

In addition to the 84 cases with confirmed molecular diagnoses and 22 with a likely molecular diagnosis, an additional 98 cases had potentially clinically significant molecular findings, which includes two heterozygous variants in a gene with autosomal recessive inheritance (two VUS or one pathogenic/likely pathogenic variant along with a VUS variant), a VUS in a gene with autosomal dominant inheritance pattern, and X-linked variants in hemizygous males classified as VUS.

The 98 cases were comprised of 56.6% cases with rod-cone dystrophy, followed by macular dystrophy, cone-rod dystrophy, cone dystrophy (Table 1). Eight cases with potentially clinically significant findings were classified based on phenotypes as ‘other’ and included cases of glaucoma, cystoid macular edema with lamellar hole, and pre-symptomatic individuals with significant family history of retinal disease.

Among these 98 cases with potentially clinically significant molecular findings, only 5 were submitted as trios. Of these 5 trio cases, 4 had two VUS in trans and one had a homozygous VUS. Eleven were submitted as duo cases. In 5 of these cases, a VUS was identified in a gene with autosomal dominant inheritance pattern and 6 cases identified two heterozygous or one homozygous variant in a gene with autosomal recessive inheritance pattern. In 4 of the 5 cases with 2 variants in a gene with autosomal recessive inheritance patterns, at least one of the identified variants met pathogenic or likely pathogenic criteria, and in 2 of these 5 cases both variants were classified as pathogenic. The remaining 83 cases were submitted as singleton proband samples.

In total, we identified variants in 85 different genes. The most common variants identified in our cohort were variants in ABC4, EYS, PRPH2, RPGGR, and USH2A, which each identified as potentially clinically significant molecular findings in more than 10 independent cases (Table 3). Variants were identified in 65.6% of cone-rod dystrophy, 64.0% of rod-cone dystrophy, 59.7% of macular dystrophy, and 30.8% of cone dystrophy cases.

### Cases with “other” phenotypes in the cohort.

Of the cases with clinical diagnoses that fell within our “Other” category, 65.4% had at least one variant identified with potential clinical significance, including 3 (11.5%) that had a molecular diagnosis. Molecular findings in this group included variants in genes such as ABC6C, CHM, GPR143, SLC38A8, ASB10, and NR2E3.

### Cases with multiple molecular findings.

In 3 cases, molecular findings in multiple genes potentially related to the clinical phenotype were identified. In one case, we reported two VUSs in two different autosomal dominant disease associated genes (Case 124; Supplemental Table 2), one case was reported with 2 heterozygous variants of unknown phase in an autosomal recessive disease gene in addition to a single VUS in an autosomal dominant disease gene (Case 93) and one case in which two heterozygous variants in three different autosomal recessive genes were identified (Case 26). Each of these cases are considered potentially clinically significant findings with a possible molecular diagnosis.

The first case (Case 124) has a VUS in two autosomal dominant genes, SNRNP200 and PRPF8. Both genes affect RNA splicing and are associated with autosomal dominant retinitis pigmentosa. Fundus examination showed numerous white dots in the inferior retina. Intraretinal pigment was noted more severely in the inferior retina than the superior retina (Supplemental Fig. 1A). ERG examination was abnormal and consistent with a
ES category | Cases approved for Luxturna | Cases enrolled/currently enrolling for intervention clinical trial | Cases provided with corrective counseling for inheritance | Cases with known family member cascade testing
--- | --- | --- | --- | ---
Molecular Diagnosis | 2 | 5 | 9 | 5
Likely Molecular Diagnosis | 1 | 0 | 1 | 0
Possible Molecular Diagnosis | 0 | 0 | 13 | 0

Table 4. Outcomes for ophthalmic disease.

Outcomes for ophthalmic disease. The ophthalmology records were reviewed to assess the outcomes of the ES testing on affected individuals and their families. The specific outcomes due to their sequencing results fell into 4 categories (1) approved for Luxturna (FDA approved gene-therapy), (2) enrolled/currently being enrolled in an interventional clinical trial, (3) correction of previously miscounselled inheritance pattern/reproductive risk, and (4) confirmed cascade testing of family members. Overall, 28 of the 357 cases (7.9%) were identified as having at least one of the four outcomes (Table 4). Of those with a molecular diagnosis 5 cases out of 84 (5.95%) were enrolled or in the process of being enrolled in an interventional clinical trial due to their sequencing results. Out of the 22 cases with a likely molecular diagnosis, 1 patient (4.5%), and her affected sibling, were able to get phase testing confirmed at a later date and have successfully completed Luxturna enrollment. ES results allowed for correction in counseling regarding inheritance and reproductive risk in 19 out of the 357 cases (5.3%). Cascade familial testing was done for 3 out 357 cases (0.8%).

In some cases variants were identified in genes causing syndromic IRD including multiple cases with biallelic variants in ciliopathy genes, Pseudoxanthoma Elasticum, ocular albinism, and Coenzyme Q10 deficiency (Supplemental Table 3). Review of the charts provided additional clinical data confirming the molecular diagnosis for some of these cases including polycystic kidneys in a patient with biallelic variants in CEP290 (Case 155), kidney failure, motor problems, and intellectual disability in a patient with a homozygous AHI1 variant (Case 221), and chronic kidney disease in a patient with two heterozygous variants of unknown phase in the NPHP4 gene (Case 16).

Secondary findings (ACMG v1 & v2). Secondary findings in the genes recommended by ACMG\textsuperscript{13,14} were reported in 4 out of 357 cases (1.12%) and included pathogenic or likely pathogenic variants in BRCA1, APOB, TNNT2, and LDLR genes.

Discussion
The extensive clinical and genetic heterogeneity of disorders of hereditary retinal degeneration have led to large scale adoption of high throughput NGS based methodologies such as gene panels, ES, and more recently whole genome sequencing (GS) which can test for a large set of genes at once. Several large NGS based studies have been published with potentially significant variants identified in 41–76% of cases, with variable yields depending on the specific disease subtype\textsuperscript{3–11} (Supplemental Table 1). The majority of these studies (6/8; 75%) utilized NGS with either targeted capture or virtual gene panel, one (12.5%) utilized ES and a stand-alone test for RPGR (ORF 15), and one (12.5%) utilized ES in a subset of probands and a targeted capture panel in the remainder of the cohort. Of these studies, only one\textsuperscript{e} used ACMG guidelines to interpret identified variants, and accordingly, in others, aspects such as the rate of confirmed molecular diagnosis versus the likely molecular diagnosis which includes two pathogenic variants with unknown phase, and whether confirmed cases also included ones with VUS or two variants with unknown phase, are unclear. The classification of variants based on ACMG guidelines and categorization of case results using a clearly defined scoring system utilized in the current study provides a consistent way to compare the diagnostic yields across different cohort datasets.

By comparison, in this study we completed clinical ES in a series of 357 individuals with inherited retinal dystrophy including specific diagnoses of rod-cone dystrophy, macular dystrophy, cone-rod dystrophy or cone dystrophy. Using ACMG guidelines to interpret all variants identified in our laboratory, we confirmed a molecular diagnosis in 23.5% of cases, a likely molecular diagnosis in an additional 6.2% of cases, and a possible molecular diagnosis in 27.4% of cases, for a total of 57.1% of cases with either a confirmed molecular diagnosis or potentially clinically significant findings. This is comparable to the yield of clinical whole exome sequencing.
Some differences between the previous studies and ours include the cohort demographics, specific clinical phenotypes, previous testing performed before inclusion in the current study, availability of familial samples for variant phasing, and NGS methodology used. The coverage and depth of reads for retinal genes may differ between targeted gene panels and ES. One of the limitations of our exome method is the inability to detect CNVs which are implicated in many retinal disorders and there are multiple studies which have underscored the importance of these variants in retinal disorders. However, unlike pediatric patients with neurodevelopmental disorders, where chromosomal microarray testing is a standard of care, eye disorder patients do not undergo standard chromosomal microarray testing, even though the literature now supports prevalence of CNVs in this cohort. CNVs along with deep intronic variants may explain a part of the missing heritability in our patient population. In at least one study, for 17.3% cases (n = 45) only a single likely causative variant in autosomal recessive cohort. CNVs along with deep intronic variants may explain a part of the missing heritability in our patient population. In at least one study, for 17.3% cases (n = 45) only a single likely causative variant in autosomal recessive disease was identified. Additional testing in a subset of these cases identified a likely causative deep intronic variant in 6 cases, a likely causative small deletion or duplication in an additional 4 cases, and 1 case in which a canonical splice variant was detected by the second testing methodology that was missed by ES likely due to low coverage. Similar to these findings, we identified a single variant in autosomal recessive disease in 21 (5.9%) cases. In 9 of these cases, single variants were identified in ABCA4, EYS, or USH2A, which all have previously been reported and/or well known pathogenic intronic variants which would not have been detected by ES. While there are few recurrent small deletions or duplications associated with some of the genes identified in these 21 samples, small private CNVs or larger CNVs cannot be excluded. Follow-up testing including GS was recommended in each of these cases, although results of those tests if conducted were not available for further analysis. The highest rate of potentially clinically significant findings as well as the highest rate of molecular diagnosis was found in individuals who were referred with a diagnosis of a genetic disorder. Individuals had a potentially clinically significant finding, and 24.6% received a molecular diagnosis. Variants in disease-associated genes were reported in 59.7% of cases with clinical findings consistent with macular dystrophy, with 22.2% of individuals receiving a molecular diagnosis. Individuals with cone dystrophy were returned potentially significant variants in 53.8% of cases, and molecular diagnoses were reported in a total of 23.1%. Interestingly, individuals with clinical findings of cone-rod dystrophy were returned the highest percentage of potentially clinically significant findings (65.6%), but molecular diagnosis was only identified in 15.6% of cases. This may reflect the small number of individuals with cone-rod dystrophy (n = 32), although it may also reflect fewer variants with significant evidence for pathogenicity.

Across all disease subtypes in our cohort, 11% of the cases had two variants identified in a disease associated with autosomal recessive inheritance with unknown phase. While follow-up testing is not always a viable option, pursuing additional testing methodologies, targeted variant testing, or segregation studies in families may ultimately resolve some of these cases. Furthermore, samples submitted to our lab were not routinely reanalyzed, so reanalysis and/or re-classification of these variants with current evidence may result in different classifications.

Limitations of our observational retrospective study include that a majority of the cases submitted were singleton samples restricting the phase information when variants in AR genes were identified, incomplete targeted familial testing for the identified variants as most of the cases were of adults, limited review of the post-ES clinical testing results which were triggered by a ES finding, and lack of systematic review on clinical follow-ups and genotype-phenotype correlation for cases with molecular findings. Further, cases analyzed before the availability of large population databases such as prior to ExAC in 2015, were not re-reviewed using the updated allele frequency information. Additionally, our clinical WES analysis approach is primarily focused on finding variants in known IRD genes and might potentially miss novel candidate gene variants.

GS is often being used on a research or clinical basis. It has the benefit of simultaneously detecting exonic variation as well as intronic variation, CNVs, and in some cases repeat expansions and mitochondrial DNA variation. While GS is currently not often utilized as a first-tier test for diagnosis of genetic disease, it has the potential to change in the near future as technology becomes more widely available and our understanding of non-exonic genetic variation in disease improves.

Methods
Retrospective review of clinical records and genetic testing results between 2012 and 2018 was performed. Patients were seen and evaluated at the Edward S. Harkness Eye Institute at Columbia University Irving Medical Center. Informed consent was waived due to the minimal risk conferred to the patients and the retrospective nature of the study design as per Columbia University Institutional Review Board-approved protocol AAAR8743. All procedures were reviewed and in accordance with the tenets of the Declaration of Helsinki. All patients underwent a full ophthalmic examination which includes assessment of best-corrected visual acuity (BCVA), acquisition of autofluorescence imaging, optical coherence tomography (OCT) scans and in most cases, full-field electroretinogram (fERG) testing. Cases with gene-specific phenotypes were sent for candidate gene sequencing. For the rest, peripheral blood samples of the proband and the family members (if available) were sent for ES testing. Since 2016, except for gene-specific phenotypes, the other cases were directly sent for ES testing.
Categorization of cases based on the phenotype. Based on the clinical examination, cases were broadly categorized into five phenotypic classifications based on the shared degenerative features and disease trajectories. The cone-rod dystrophy classification encompasses disorders characterized by initial pan-retinal rod degeneration with subsequent cone involvement typified by conditions such as LCA, retinitis pigmentosa and related disorders. The cone-rod dystrophy classification includes disorders involving selective degeneration of cones such as achromatopsia and macular dystrophies include degenerative disorders confined to the posterior pole such as Stargardt disease or pattern macular dystrophy. The “other” category included cases of myopic degeneration, choroideremia, ocular albinism, glaucoma, familial drusen, pseudo-xanthoma elasticum, etc.

Clinical exome sequencing. For clinical ES, libraries were prepared from genomic DNA from the proband and the parents using Agilent SureSelectXT (Human All Exon v5 + UTRs) capture kit according to the manufacturers’ protocol. Paired-end sequencing was performed on the Illumina HiSeq 2500 platform (average coverage of 100x). The sequence data was aligned [human genome reference sequence (GRCh37/hg19)] and annotated using Nextgene (version 2.3; Softgenetics, LLC. PA) software. Variant filtering and annotation were performed using an in-house developed pipeline and reviewed as part of the clinical workflow for constitutional clinical ES. The WES variants were categorized into various bins: (1) All variants that had been reported as pathogenic, (2) All variants in genes with associated phenotypes, and (3) All variants in genes not associated with phenotypes. These three bins were then filtered at 1% population frequency using 1000 genomes, Exome Variant Server\(^\text{18}\) and were used to generate a total of six separate outputs. Each of these output had variants in different tabs: disruptive variants (frameshift, stopgain, canonical splice site), missense variants, two or more heterozygous variants per gene (potential compound heterozygous variants), De Novo variants (for trios), Unique variants (variants only present in the proband and not seen in our database) and homozgyous/hemizygous variants. Additional population databases such as ExAC and gnomAD were used for variant prioritization as they became available.

Identified variants were assessed for the phenotype match by means of a bi-weekly case discussion meeting with the clinical team. Variants were classified using the guidelines from the ACMG and the Association of Molecular Pathology (AMP)\(^\text{12}\); Variants classified as pathogenic, likely pathogenic and VUS were reported in the test results. Heterozygous pathogenic/likely pathogenic variants in autosomal recessive disease associated genes (carriers) were also reported with a recommendation for a reflex to Genome Sequencing (GS) or copy number variant (CNV) test at an outside provider since these tests are currently not performed at our laboratory. Routine re-analysis of ES data is currently not part of our constitutional case analysis workflow in the clinical laboratory, hence other than special instances most cases have not been re-analyzed.

RPGR ORF15 variants: amplicon based sequencing. RPGR ORF15 testing was performed on selected samples, which were negative by clinical ES test and met clinical indication for RPGR ORF15 testing. RPGR ORF15 variant detection was performed using a long range PCR, amplicon based NGS and analysis method. For this testing the genomic region chrX:38144742-38146527 (hg19) corresponding to the RPGR ORF15 region was amplified by PCR using the following primers, ORF15F 5’-GTA TGA TTT TAA ATG TGA TCG CTT GTC AGA and ORF15R 5’-GAAGGGATTAAAAATGTGATCGCTTGCAGA G and ORF15R 5’-AA-AGGGACATTTAAAAATGTGACTGCTG GCCATAATC\(^\text{19}\). PCR was performed using standard methods using Platinum SuperFi PCR Master Mix (Invitrogen Cat.# 12358-010). The 1.7 kb PCR product was fragmented into smaller sized DNA, typically 150–200 bp, purified and end-repaired following 3’adenylation. Double stranded sequencing adaptors were ligated to both ends of this DNA. The library was loaded onto the MiSeq instrument (Illumina) for cluster generation and sequencing. Analysis of the sequence reads was performed by NextGene software (SoftGenetics, LLC. Pennsylvania). For data analysis, the “consolidation” tool in NextGENE software was used, which merges overlapping sequences to give a consensus sequence in place of all the original reads that are in the subregion. This method is recommended for datasets that have a high depth of coverage in the raw reads, when there is a need to correct sequence reads such as those in repeat regions and difficult to align repeat rich regions. The resulting variant calls from the original reads and the consolidation output were evaluated for validity, based on variant allele fractions, gender of the proband and if the variant was seen in ES reads from the ORF15 region.

Categorization of cases based on the results of ES test. Based on the ES results, the laboratory classified the cases into four categories. (1) Cases in which a pathogenic or likely pathogenic variant in an autosomal dominant (AD) disease gene, an X-Linked (XL) pathogenic or likely pathogenic variant in hemizygous or homozygous state, or in cases in which two pathogenic or likely pathogenic variants were identified in autosomal recessive (AR) disease in trans, in a disease gene with clinical phenotype match were classified as a “Molecular Diagnosis”. (2) Cases in which two pathogenic or likely pathogenic variants explaining the phenotype were identified with unknown phase were categorized as “Likely Molecular Diagnosis”. (3) Cases with two heterogeneous variants with unknown phase in an AR disease gene (two VUS or one pathogenic/likely pathogenic variant along with a VUS variant), or cases in which a VUS was identified in an AD or XL gene were categorized as “Possible Molecular Diagnosis” (4) Remaining cases in which no variants potentially explaining the clinical phenotype were identified were categorized as “Negative”.

Data availability All variants reported during this study are included in this published article including supplementary information files. The raw next-generation sequencing data used to identify the variants reported herein cannot be made public due to the sensitive nature of the data. However, datasets generated and analyzed during the current study are available in the NCBI BioProject repository with the accession number PRJNA749803.
publicly available for reasons of patient confidentiality and in compliance with HIPAA regulations. Qualified researchers or clinicians may apply for access to these data pending institutional review board approval.

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References
1. Russell, S. et al. Efficacy and safety of voretigene neapavovec (AAV2-hRPE65v2) in patients with RPE65-mediated inherited retinal dystrophy: A randomised, controlled, open-label, phase 3 trial. Lancet 390, 849–860 (2017).
2. Miyadera, K. et al. Targeting ON-bipolar cells by AAV gene therapy stably reverses LRIT3-congenital stationary night blindness. Proc. Natl. Acad. Sci. USA 119, e2117038119 (2022).
3. Wang, P. et al. An ophthalmic targeted exome sequencing panel as a powerful tool to identify causative mutations in patients suspected of hereditary eye diseases. Transl. Vis. Sci. Technol. 8, 21 (2019).
4. Carss, K. J. et al. Comprehensive rare variant analysis via whole-genome sequencing to determine the molecular pathology of inherited retinal disease. Am. J. Hum. Genet. 100, 75–90 (2017).
5. Haer-Wigman, L. et al. Diagnostic exome sequencing in 266 Dutch patients with visual impairment. Eur. J. Hum. Genet. 25, 591–599 (2017).
6. Wang, L. et al. Application of whole exome and targeted panel sequencing in the clinical molecular diagnosis of 319 Chinese families with inherited retinal dystrophy and comparison study. Genes (Basel) 9, 360 (2018).
7. Huang, X. F. et al. Genotype-phenotype correlation and mutation spectrum in a large cohort of patients with inherited retinal dystrophy revealed by next-generation sequencing. Genet. Med. 17, 271–278 (2015).
8. Stone, E. M. et al. Clinically focused molecular investigation of 1000 consecutive families with inherited retinal disease. Ophthalmology 124, 1314–1331 (2017).
9. Ellingford, J. M. et al. Molecular findings from 537 individuals with inherited retinal disease. J. Med. Genet. 53, 761–767 (2016).
10. Carrigan, M. et al. Panel-based population next-generation sequencing for inherited retinal degenerations. Sci. Rep. 6, 33248 (2016).
11. Agnelli, L. et al. Next-generation sequencing applications for inherited retinal diseases. Int. J. Mol. Sci. 22, 5684 (2021).
12. Richards, S. et al. Standards and guidelines for the interpretation of sequence variants: A joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. Genet. Med. 17, 405–424 (2015).
13. Green, R. C. et al. ACMG recommendations for reporting of incidental findings in clinical exome and genome sequencing. Genet. Med. 15, 365–574 (2013).
14. Kalia, S. S. et al. Recommendations for reporting of secondary findings in clinical exome and genome sequencing, 2016 update (ACMG SF v2.0): a policy statement of the American College of Medical Genetics and Genomics. Genet. Med. 19, 249–255 (2017).
15. Yang, Y. et al. Clinical whole-exome sequencing for the diagnosis of mendelian disorders. N. Engl. J. Med. 369, 1502–1511 (2013).
16. Ellingford, J. M. et al. Assessment of the incorporation of CNV surveillance into gene panel next-generation sequencing testing for inherited retinal diseases. J. Med. Genet. 55, 114–121 (2018).
17. Van Schil, K. et al. Mapping the genomic landscape of inherited retinal disease genes prioritizes genes prone to coding and non-coding copy-number variations. Genet. Med. 20, 202–213 (2018).
18. Wang, Y. et al. A mutation abolishing the ZMPSTE24 cleavage site in prelamin A causes a progeroid disorder. J. Cell Sci. 129, 1975–1980 (2016).
19. Ruddle, J. et al. RPGR OF15 genotype and clinical variability of retinal degeneration in an Australian population. Br. J. Ophthalmol. 93, 1151–1154 (2009).

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Competing interests
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