A combined RNA-seq and whole genome

² sequencing approach for identification of

non-coding pathogenic variants in single

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19 Abstract

20	Inherited retinal degenerations (IRDs) are at the focus of current genetic
21	therapeutic advancements. For a genetic treatment such as gene therapy to be
22	successful an accurate genetic diagnostic is required. Genetic diagnostics relies
23	on the assessment of the probability that a given DNA variant is pathogenic.
24	Non-coding variants present a unique challenge for such assessments as
25	compared to coding variants. For one, non-coding variants are present at much
26	higher number in the genome than coding variants. In addition, our
27	understanding of the rules that govern the non-coding regions of the genome is
28	less complete than our understanding of the coding regions. Methods that allow
29	for both the identification of candidate non-coding pathogenic variants and their
30	functional validation may help overcome these caveats allowing for a greater
31	number of patients to benefit from advancements in genetic therapeutics. We
32	present here an unbiased approach combining whole genome sequencing
33	(WGS) with patient induced pluripotent stem cell (iPSC) derived retinal organoids
34	(ROs) transcriptome analysis. With this approach we identified and functionally
35	validated a novel pathogenic non-coding variant in a small family with a
36	previously unresolved genetic diagnosis.

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38 Introduction

39 Inherited retinal degenerations (IRDs) are a leading cause of blindness, 40 altogether affecting >2million people worldwide. IRDs are characterized by progressive degeneration of photoreceptor and/or retinal pigment epithelial 41 (RPE) cells of the retina with variable age of onset and rates of degeneration¹. 42 43 Despite tremendous ongoing efforts and research into various therapeutics, treatments for IRDs remain limited. Currently there are two regulatory agency-44 approved treatment approaches, retinal prosthesis implants and gene 45 augmentation therapy for IRD caused by mutations in the RPE65 gene $^{1-7}$. The 46 eye is a prime candidate for gene therapy approaches due to its relative immune-47 privilege, surgical accessibility and ease of non-invasive monitoring. In addition, 48 IRDs are Mendelian disorders caused by mutations in single genes in the vast 49 majority of cases. Owing to these favorable circumstances several gene/genetic 50 51 therapy clinical trials have been initiated for IRDs, including those caused by mutations in the ABCA4, CEP290, CHM, CNGA3, CNGB3, MYO7A, RPGR, 52 RS1, and USH2A⁶. 53

As each genetic therapy targets a specific gene, for a patient to be considered for treatment they must obtain a reliable genetic diagnosis. Difficulties inherent to genetic diagnostics are rooted in the fact that every individual carries millions of DNA variants in their genome ^{8,9}. The large majority of the DNA variants are found in non-coding regions of the genome such as intergenic and intronic regions. Since non-coding sequences can better tolerate sequence variation

60	compared to coding sequences, most of these variants are benign and do not
61	lead to disease. Still, some non-coding variants are found to be pathogenic by
62	altering gene expression and/or splicing patterns ^{10–12} . As a result, non-coding
63	variants are among the hardest to classify and thus under-diagnosed ^{13,14} .
64	Algorithms exist that predict the effect of a non-coding variant on gene
65	expression or splicing based on analysis of the DNA sequence alone, but their
66	accuracy for diagnostic purposes remains undetermined ^{15–18} .
67	In order to functionally test the effects of non-coding variants one needs to
68	quantify the level of gene expression and analyze the splicing patterns of the
69	presumably affected genes. When multiple variants in multiple genes need to be
70	evaluated, advanced methods for whole transcriptome analysis are
71	advantageous. Indeed, previous studies successfully utilized large RNA-seq
72	datasets from tissue biopsies to identify novel non-coding pathogenic variants
73	^{19,20} . For example, Evrony <i>et al</i> narrowed down a linkage analysis in a very large
74	pedigree to a single non-coding variant using RNA-seq. This non-coding variant
75	was shown to cause intron retention in the DONSON gene and is most likely the
76	genetic cause of microcephaly-micromelia syndrome (MMS) in this population ¹⁹ .
77	Cummings et al used 184 skeletal muscle RNA-seq samples available through
78	Genotype-Tissue Expression resource (GTEx) ²¹ as a reference panel for 50
79	patients with undiagnosed muscle disorders. This comparison led to a genetic
80	diagnosis for 17 previously unsolved families and identification of several splice
81	altering variants ²⁰ .

82 Such studies depend on the availability of large, publicly available RNA-seq datasets and/or on a large cohort of patients. They also require biopsy samples 83 from a clinically relevant tissue or cell type. Both gene expression and splicing 84 are tissue-specific ^{22,23} owing to restricted availability of transcription and splicing 85 factors with variable usage of regulatory DNA sequences ^{24–27}. Consequently, 86 DNA variants in such regulatory sequences can have tissue-specific outcomes 87 on gene expression and splicing ^{18,28}. Thus, analyzing RNA from a clinically 88 relevant tissue or cell type is crucial to obtain a more focused and reliable 89 90 diagnostic result. When a clinically relevant tissue is not accessible, ex vivo surrogate models can sometimes suffice. Indeed, a study aimed at discerning the 91 genetic causality of patients with monogenetic neuromuscular disorders found 92 that t-myotubes, skeletal myotubes derived by myoD overexpression in 93 fibroblasts, accurately reflected the muscle transcriptome and faithfully revealed 94 pathogenic variants ²⁹. 95

In this study, we aimed to develop a pipeline that would detect putative non-96 coding pathogenic mutations in a small family. We present a pilot study 97 performed in a five member family with two siblings affected by cone dysfunction 98 syndrome in which we successfully identify and functionally validate a novel deep 99 100 intronic variant without the use of large reference datasets. Obtaining a clinically relevant tissue from IRD patients is not possible as the retina cannot be safely 101 102 biopsied. To overcome this limitation, we have made use of patient-derived induced pluripotent stem cells (iPSCs) that were differentiated in vitro to form 103 retinal organoids (ROs)³⁰. ROs have been shown to recapitulate many aspects of 104

human retinal structure and function ^{30,31,40,32–39}. We show that the RO
transcriptome is much closer to the transcriptome of normal human retina than
other more readily available diagnostic tissues. More importantly, we show for
the first time that analysis of a patient-derived RO transcriptome can successfully
detect pathogenic deep intronic variants that activate cryptic splice sites, leading
to a new genetic diagnosis. Our approach can lead to a larger number of patients
to be eligible for genetic therapies.

112 Materials and Methods

113 Human Subjects

114 The study was approved by the institutional review board at the Massachusetts

115 Eye and Ear (Human Studies Committee MEE in USA) and adhered to the

116 Declaration of Helsinki. Informed consent was obtained from all individuals on

117 whom genetic testing and further molecular evaluations were performed.

118 Pluripotent Stem Cell Induction

Tissue samples were obtained with written informed consent in adherence with 119 the Declaration of Helsinki and with approval from institutional review boards at 120 the University of Wisconsin-Madison and Massachusetts Eye and Ear Infirmary. 121 Blood samples from 4 individuals from family OGI-081 (197, 198, 200 and 340) 122 were collected and reprogrammed by Cellular Dynamics, Inc (now FUJIFILM 123 Cellular Dynamics, Inc) as custom MyCell products. Three independent clones 124 from each individual were karyotypically normal, expressed pluripotency markers 125 and successfully differentiated to retinal organoids (⁴¹: lines 1579, 1580, 1581 126

- and 1582). Stem cells were maintained on Matrigel (ThermoFisher) in either
- 128 mTeSR1 (WiCell) or Stemflex (ThermoFisher) and passaged with either Versene
- 129 or ReLeSR (STEMCELL Technologies).
- 130 Retinal Organoid (RO) Differentiation
- 131 Differentiation of iPSCs was performed as previously described ⁴¹. Briefly,
- embryoid bodies (EB) were lifted with either 2 mg/ml dispase or ReLeSR and
- weaned into Neural Induction Media (NIM: DMEM:F12 1:1, 1% N2 supplement,
- 134 1x MEM nonessential amino acids (MEM NEAA), 1x GlutaMAX and 2 mg/ml
- heparin (Sigma)) over the course of 4 days. On day 6, 1.5 nM BMP4 (R&D
- 136 Systems) was added to fresh NIM and on day 7, EBs were plated on Matrigel at
- a density of 200 EBs per well of a 6-well plate. Half the media was replaced with
- 138 fresh NIM on days 9, 12 and 15 to gradually dilute the BMP4 and on day16, the
- media was changed to Retinal Differentiation Media (RDM: DMEM:F12 3:1, 2%
- B27 supplement, MEM NEAA, 1X antibiotic, anti-mycotic and 1x GlutaMAX). On
- 141 days 25-30, optic vesicle-like structures were manually dissected and maintained
- as free floating organoids in poly HEMA (Sigma)- coated flasks with twice weekly
- feeding of 3D-RDM (RDM + 5% FBS (WiCell), 100 μ M taurine (Sigma) and
- 144 1:1000 chemically defined lipid supplement) to which 1 μM all-trans retinoic acid
- 145 (Sigma) was added until d100. Live cultures were imaged on a Nikon Ts2-FL
- 146 equipped with a DS-fi3 camera.
- 147 Immunocytochemistry and Microscopy

148 Organoids were fixed in 4% paraformaldehyde at room temperature for 40 min,

cryopreserved in 15% sucrose followed by equilibration in 30% sucrose, and

150	sectioned on a cryostat. Sections were blocked for 1 hr at room temperature (RT)
151	in 10% normal donkey serum, 5% BSA, 1% fish gelatin and 0.5% Triton then
152	incubated overnight at 4° C with primary antibodies diluted in block. Table S1 lists
153	primary antibodies, dilutions and sources. Slides were incubated with species-
154	specific fluorophore-conjugated secondary antibodies diluted 1:500 in block, for
155	30 minutes in the dark at RT (Alexa Fluor 488, AF546 and AF647). Sections
156	were imaged on a Nikon A1R-HD laser scanning confocal microscope (Nikon
157	Corporation, Tokyo, Japan).
158	DNA Sequencing
159	DNA was extracted from venous blood using the DNeasy Blood and Tissue Kit
160	(Qiagen, Hilden, Germany). OGI-081-197 underwent GEDi sequencing as
161	described previously ⁴² . All five family members underwent whole exome and
162	PCR-free whole genome sequencing. Sequencing was done at the Genomics
163	Core at Massachussets Eye and Ear as described previously ⁴³ .
164	RNA Sequencing
165	For transcriptome analysis, ROs from at least 2 different clones per individual
166	were harvested at approximately day160 (early stage 3), lysed in 350 μ l buffer
167	RLT+ME from the RNeasy mini kit (Qiagen), snap frozen on dry ice and stored at
168	-80C. At a later time samples were defrosted on ice and passed through
169	QIAshredder columns (Qiagen). Subsequently, Total RNA was extracted per the
170	manufacturer's instructions. RNA quality and quantity was assessed on an
171	Agilent 2100 Bioanalyzer, RIN number ranged between 9.6-9.9. For each sample
172	$1\mu g$ of total RNA spiked with 1.2ng Sequins (v2) controls 44 was used to generate

173 RNA-seq paired-end libraries with the Illumina TruSeq Stranded Total RNA kit.

174 Ribosomal RNA was removed with the Ribo-Zero Human/Mouse/Rat kit.

Libraries were multiplexed and sequenced on an Illumina HiSeq 2500 instrument

176 for 101 cycles.

177 Bioinformatics

178 Whole exome sequence data was analyzed in house ⁴³ and whole genome data

was analyzed in collaboration with the Broad Institute of MIT and Harvard using

180 methodology described previously ²⁰. Briefly, BWA was used for alignment.

181 GATK was used for single nucleotide polymorphism and insertion/deletion calls.

182 Additional variant annotation was performed using the Variant Effect Predictor

(VEP) ⁴⁵. Variants of interest were limited to polymorphisms with less than 0.005

allelic frequency in the gnomAD and ExAC databases ^{8,9}. Whole genome copy

number analysis, with consideration of structural changes, was done using
 Genome STRiP 2.0 ⁴⁶.

187 For analyses of RNA-seq data, read quality was assessed with FastQC v0.11.3

(Babraham Bioinformatics, Cambridge, UK) and MultiQC v1.2⁴⁷. Reads were

aligned to the human genome version GRCh37 by the STAR v2.5.3a ⁴⁸ aligner in

190 two-pass mode within the sample and across replicates for each sample sets.

191 Annotations were derived from the Human GENCODE v19 (Ensemble74).

FeatureCount v1.5.2 ⁴⁹ from the Subread package, was used to generate gene expression matrix with the following non-default settings, reads must be paired, both the pairs must be mapped, use only uniquely mapped reads, multi-mapped reads are not counted, chimeric reads are not counted and strand specificity

turned on. Anaguin ⁵⁰ was further used to evaluate alignment sensitivity and gene 196 expression. Here sensitivity indicates the fraction of annotated regions covered 197 by alignments of the reads by STAR (Table S2). No limit of quantification or limit 198 of detection was reported. 199 200 For discovery of novel or known alternative splicing events we used a combination of CASH v2.2.1 ⁵¹ and MAJIQ v1.1.3a ⁵². CASH was operated with 201 default settings. MAJIQ was run with a minimum of 5 reads for junction detection 202 and 10 reads for the calculation of delta percent spliced-in (dPSI). EdgeR 203 (v3.2.2) ⁵³ was used to perform differential gene expression with default settings. 204 205 Data normalization was performed using trimmed mean of M-values (TMM). Next, the raw read counts were converted to transcript per million (TPM) 206 expression values. The Picard tools v1.87 and RSeQC v2.6.4⁵⁴ were used to 207 calculate mean fragment length. The approach implemented in Kallisto ⁵⁵ was 208 used to covert raw reads to TPM values. An average TPM of the third lowest 209 Sequins between test and control samples was calculated and used as cutoff. 210 TPM values for the GTEx samples used in figure3 were downloaded from the 211 GTEx portal. The human normal retina (HNR) samples ⁵⁶ and the ENCODE skin 212 213 samples were reanalyzed as described above. The ENCODE skin samples were used for the analysis performed with MAJIQ as they were generated with a 214 215 stranded total RNA-seq library same as our RO samples. In contrast the GTEx 216 samples were generated with an mRNA non-stranded library. 217

218 RT-PCR and cloning

- 219 RT-PCR was conducted using SuperScript IV first-Strand synthesis system
- (ThermoFisher Scientific, Waltham, MA). Exon 14b was amplified with primers F:
- 221 GACATGTTGCTAAGATTGAAATCCGT from exon 14 and R:
- 222 GACCCAGCTTTCAGAGTAACCAGAAC from exon 15 using Phusion
- polymerase (NEB, Ipswich, MA). The longer band containing exon14b was then
- excised from the gel and purified using Zymoclean gel DNA recovery kit
- 225 (Zymoresearch, Irvine, CA) and cloned using pGEM-T Easy Vector System
- (Promega, Madison, WI). The plasmid was used in a transformation into
- 227 Subcloning Efficiency DH5α Competent Cells (Invitrogen, Carlsbad, CA). The
- 228 plasmid was isolated with Zyppy Plasmid Miniprep Kit (Zymoresearch, Irvine,
- 229 CA). All procedures described in this section were conducted according to the
- 230 manufacturer instructions. The DNA sequence of Exon 14b was found to be:
- 231 GCCAGGTGCAGTGGCTCACGACTGTAATTCCAACACTTTGGGAGGCCAAGG
- 232 TGGCAGGATCACATAAGTCCAGGAGTTCAAGACAAGCCTGGACAACATG.

234 **Results**

235 <u>Unresolved genetic analysis of family OGI-081</u>

The pilot study reported here was conducted on a five member family with two 236 237 siblings shown by clinical testing to be affected by a cone dysfunction syndrome (Figure1a). Both affected patients had nystagmus and decreased vision from 238 239 infancy and at age 8, OGI-081-197 was also noted to have photophobia. Visual 240 acuity for both affected patients measured 20/150-200 and remained stable for 3 241 years. Full field electroretinogram (ffERG) testing of retinal function for OGI-081-197 was significant for reduced and delayed cone photoreceptor responses, with 242 243 normal rod photoreceptor response amplitudes. Optical coherence tomography (OCT) imaging of the retina showed retinal degenerative changes in the fovea 244 (Figure1b). In addition to their retinal disease, both affected patients were found 245 to have Chiari malformations. Interestingly, vision phenotypes such as 246 photophobia, vision loss and nystagmus have been reported as accompanying 247 symptoms in some forms of Chiari malformations^{57–60}. Unfortunately, despite 248 evidence that Chiari malformations have a hereditable component ^{61–63}, the 249 genes involved are not yet well defined ^{59,64,65}. For that reason we could not rule 250 251 out the possibility that the vision phenotypes and Chiari malformations share a common genetic causality. 252

The phenotypes segregation in OGI-081 is indicative of a recessive mode of inheritance. Thus, for genetic testing we searched for genes that have putatively pathogenic variants in both alleles (Figure1a). Selective exon capture based

256 genetic diagnostic testing was performed using the Genetic Eye Disease (GEDi) test ⁴². Since this did not identify a clear cause of disease, whole exome 257 sequencing (WES) for the five members of the family was performed. Both the 258 GEDi testing and WES identified a single rare variant in the CNGB3 gene 259 (c.1148delC, p.Thr383llefsTer13) which has been reported to be pathogenic ^{66–68}. 260 but a second rare variant in CNGB3 was not identified, nor were other potential 261 causative genetic variants forthcoming for the two affected members of the 262 family. CNGB3 mutations are among the most common causes of cone 263 264 dysfunction syndrome, but to the best of our knowledge, Chiari malformations have not been reported as an accompanying symptom in CNGB3 patients ^{69,70}. 265 Moreover, the 1.75e⁻³ gnomAD allele frequency of the p.Thr383llefsTer13 variant 266 is higher than expected for recessive pathogenic variants and two homozygous 267 individuals are reported in the gnomAD database. Since it has been proposed 268 that up to 1 in 4-5 individuals in the general population may be a carrier of null 269 mutations in IRD genes ⁷¹ it was possible that the presence of variant 270 p.Thr383llefsTer13 was an incidental finding. 271

We therefore decided to test for two possible disease scenarios. One is that *CNGB3* accounts for the cone dysfunction syndrome and the second allele is a
non-coding variant. In this case the Chiari malformation has a separate,
unrelated causality. The second possibility is that a novel disease-causing gene
is responsible for both the cone dysfunction syndrome and the Chiari
malformation. To test these hypotheses, we performed both whole genome
sequencing (WGS) and RNA-seq analysis of a surrogate retinal tissue to

determine whether the combination of these orthogonal investigations could yielda clear genetic solution.

281	Analysis of DNA variants detected by WGS identified 3268 segregating rare
282	variants that could be sorted into 8191 allelic pairs in 642 genes (TableS3). The
283	variant ranked at the top of the list of potential causes of disease remained the
284	known pathogenic variant c.1148delC; p.Thr383llefsTer13 in the CNGB3 gene.
285	However, a second coding variant once again was not found in this gene. Next,
286	we set to establish a clinically relevant surrogate transcriptome for the human
287	retina.

288 The iPSC-derived retinal organoid (RO) transcriptome can be used as a

289 <u>surrogate for a human retinal biopsy</u>

Patient-derived iPSCs were generated from peripheral blood monocytes of all 290 members of family OGI-081 excluding OGI-081-199. The iPSCs were subjected 291 to an *in vitro* differentiation process to generate ROs with attached RPE (Figure 292 2a) ³⁰. RPE was specifically retained in the ROs so as to concurrently identify 293 potential mutations in RPE genes as well as photoreceptor genes (Figure2b). 294 The ROs were kept in culture for 160 days (early stage 3³⁰), a time point at 295 which outer segments are visible by light microscopy and cone and rod cells are 296 clearly distinguished by immunocytochemistry (Figure 2b, c-h). At da160, ROs 297 298 were harvested for total RNA isolation, library preparation, and RNA-seg analysis (Table1). 299

300 In order to evaluate if day 160 ROs can be used as informative surrogates for the adult human retinal transcriptome, we examined the expression levels of 270 301 known IRD genes reported in the RetNet database. For this analysis, we 302 compared an in-house dataset composed of 3 post-mortem human normal 303 retinas (HNR, N=3)⁵⁶ to ROs derived from the unaffected sibling (N=5) or skin or 304 whole blood samples taken from the GTEx database (Figure 3 and Table S4) 21 . 305 Skin and blood represent tissues that are more readily available - and thus 306 commonly used – for surrogate diagnostic testing. We found 224 IRD genes to 307 308 be expressed in HNR (TPM>1). Interestingly we were able to detect expression of 254 IRD disease genes in the RO samples (Figure 3 and TableS4). The higher 309 detection rate of disease-causing genes in ROs compared to HNR is most likely 310 because of overall higher TPM values in RO samples (Figure 3a&b), possibly 311 due to higher RNA quality compared to post-mortem HNR samples. In addition, 312 the presence of RPE and photoreceptors with varied maturation statuses in the 313 RO samples could be contributing factors to this finding. As expected, skin and 314 blood expressed lower numbers of IRD genes (188 & 130 respectively) at much 315 316 lower TPM values (Figure 3a&b and Table S4). Since IRD genes were very poorly represented in the blood samples, we excluded blood from further 317 318 analysis.

We next examined the complexity of the HNR, RO and skin transcriptomes, which is reflected by the multitude of isoforms that are produced from each gene locus ^{72,73}. Isoform diversity that occurs via alternative splicing of the pre-mRNA can be represented by the splice junctions detected in each gene locus. We used

the MAJIQ ⁵² algorithm to detect splice junctions in HNR and RO samples of the 323 OGI-081 unaffected sibling, and the ENCODE database ^{74,75} to detect splice 324 junctions in corresponding skin samples. The skin samples from the ENCODE 325 database were more suited for this analysis then the GTEx samples due to RNA-326 seq library type (see Materials and Methods). We found a comparable number of 327 splice junctions in the HNR and RO samples (41,121 and 31,535 (76%) 328 respectively) but a much lower number in the skin samples (16,713 (40%)) 329 (Figure 3c and TableS5). This result is probably due to the fact that HNR and 330 331 ROs are composed of a more diverse cell population than skin. More importantly, in IRD genes, the gap in complexity between the HNR (946 junctions) and ROs 332 (739 (78%)) as compared to skin (227(24%)) is even greater (Figure 3d & Table 333 S5). Thus, the RO transcriptome provided a close facsimile of the human retinal 334 and IRD transcriptomes at the gene expression and splicing pattern levels, 335 336 whereas the skin transcriptome did not.

337 Detection of a novel non-coding pathogenic variant in CNGB3

In order to find the underlying genetic cause of the retinal degeneration in OGI-338 081 affected patients, we conducted differential splicing and gene expression 339 340 analyses of the RNA-seg data obtained from day 160 ROs of affected versus 341 unaffected siblings. The differential splicing analysis was conducted with CASH ⁵¹ and MAJIQ ⁵² algorithms, and we used edgeR algorithm ⁵³ for differential gene 342 expression. CASH detected 106 differential splicing events in 101 genes (Table 343 344 S6), while MAJIQ detected 522 differential splicing junctions in 260 genes (Table S7). A comparison to the 642 genes with DNA variant pairs indicated that only 345

two genes, *CNGB3* and *NCALD*, had altered splicing patterns and a DNA variant
in each of their alleles that segregated according to disease status in OGI-081
(Figure 4a).

For the NCALD gene, the alternative splicing events identified by CASH and 349 350 MAJIQ were isoform switching events between minor isoforms (data not shown) whose biological significances are unclear. In addition, the DNA variants in this 351 352 gene are located 15-50kb away from the nearest alternative junction (Figure 4b 353 and Table S8). Given such large distances, it is not likely that these variants could cause the alternative splicing events. Moreover, the differential gene 354 355 expression analysis did not find the NCALD gene to be differentially expressed between the affected and unaffected siblings (Table S9). Taken together, these 356 357 results indicate that the NCALD gene variants are most likely not the genetic cause for the disease phenotypes observed in OGI-081. 358 359 We next examined the CNGB3 gene locus. Based on segregation in the family, we found one allele carrying the intronic variant chr8:g.87618576G>A while the 360 second allele carried the known pathogenic variant c.1148delC; 361 p.Thr383llefsTer13, and a second intronic variant chr8:g.87676221T>C (Table 362 S3). Both MAJIQ and CASH detected an alternative splicing event spanning 363 364 variant chr8:g.87618576G>A. In contrast, no alternative splicing events were 365 found to span variant chr8:g.87676221T>C (Figure4b and TableS6-8). Close examination of the alternative splicing event spanning variant 366 367 chr8:g.87618576G>A revealed that it incorporated a cryptic exon into CNGB3 in RNA samples taken from both affected siblings but not from the unaffected 368

369	sibling (Figure 5a&b). The cryptic exon is spliced between canonical exon 14 and
370	exon 15 and will therefore be termed exon14b from hereon. The inclusion of
371	exon 14b was validated by RT-PCR and subsequent cloning and Sanger
372	sequencing of the novel longer isoform from the two affected siblings (Figure 5b).
373	Both the addition of exon 14b to the CNGB3 transcript as a result of variant
374	chr8:g.87618576G>A, and the single base pair deletion in the second allele
375	carrying variant c.1148delC; p.Thr383llefsTer13, lead to a frame shift and
376	subsequent premature termination. We therefore expected both alleles to
377	undergo nonsense mediated decay (NMD) with down regulation of CNGB3
378	mRNA levels in the affected siblings as compared to the unaffected sibling.
379	However, contrary to our expectations, CNGB3 expression was not significantly
380	down regulated in our RNA-seq dataset, as analyzed using the edgeR program.
381	A comparison of each of the affected siblings with the unaffected sibling yielded
382	log_2FC values of -1.05 and -1.13, indicating slightly lower expression levels in the
383	affected siblings that did not reach statistical significance (p-values of 0.13 and
384	0.16 respectively (Table S9)). These two frame shift alleles are predicted to
385	encode truncated proteins. The protein encoded by the exon 14b including
386	isoform is predicted to maintaining a full transmembrane domain but lack the
387	ligand binding domain of CNGB3 (Figure 5c). Similarly, the protein encoded by
388	the isoform carrying the known pathogenic variant c.1148delC;
389	p.Thr383llefsTer13 is predicted to have a truncated transmembrane domain in
390	addition to lacking the ligand binding domain (Figure 5c). In order to determine
391	whether the truncated CNGB3 proteins are being translated, we performed

392 immunohistochemistry on ROs from the exon 14b allele carrier parent (OGI-081-200) and one affected sibling (OGI-081-197, Figure 6). CNGB3 is a subunit of the 393 394 cone cyclic nucleotide-gated (CNG) channel, which localizes to cone photoreceptor outer segments in chicken and mice ^{76,77}. We have also validated 395 the localization of human CNGB3 to cone photoreceptor outer segments in the 396 397 human retina (Figure S2). We therefore immunostained ROs for CNGB3 and ML opsin, the latter serving as a marker for photoreceptor outer segments. For these 398 399 studies, stage 3 ROs were kept in culture for a total of 262 days, allowing cones 400 full opportunity to mature and localize ML opsin and CNGB3 to the photoreceptor outer segments. As expected, CNGB3 co-localized with ML opsin in cone 401 photoreceptor outer segments in the parent (Figure 6c), with weaker staining 402 observed in inner segments as well (Figure 6b&c), presumably due to 403 404 mislocalization of truncated CNGB3 protein produced by the exon 14b including 405 allele. In ROs from the affected sibling, where both alleles are predicted to result in truncated proteins, CNGB3 was only observed diffusely in the cell body and in 406 inner segments; i.e., no co-localization with ML opsin was observed in cone 407 408 photoreceptor outer segments (Figure 6e&f). Taken together, results from the differential splicing analysis indicate that the likely cause for the inherited retinal 409 410 degeneration in OGI-081 is two pathogenic alleles in CNGB3 - the known pathogenic allele p.Thr383llefsTer13 and the novel deep intronic allele 411 412 chr8:g.87618576G>A; NM_019098.3:c.1663 – 2137C>T; pLeu524 llefsTer50.

413

414 Splicing prediction algorithms

415	With the identification of the non-coding pathogenic variant in CNGB3, we set out
416	to examine the mechanism by which it promotes the inclusion of exon 14b. We
417	analyzed the splice junctions surrounding exon14b with the variant analysis tool
418	Alamut Visual. Alamut Visual incorporates three splicing predictors capable of
419	analyzing deep intronic variants, SpliceSiteFinder-like (SSF) ⁷⁸ , MaxEntScan ⁷⁹
420	and NNSPLICE ⁸⁰ . All three algorithms predicted chr8:g.87618576G>A to
421	strengthen a cryptic donor splice site (DSS) (Table 2). All three algorithms also
422	detected a potential acceptor splice site (ASS) at position c.1663-2238, exactly
423	where our Sanger sequencing indicated the acceptor site of exon 14b resides.
424	Interestingly, exon 14b ASS is a stronger than the one located at exon 15 (Table
425	2). It is plausible that the availability of this acceptor site and its ability to compete
426	with the acceptor site of exon 15 contributed to the effect of variant
427	chr8:g.87618576G>A on the splicing pattern of CNGB3 in the affected siblings. In
428	addition we noticed the presence of a second even stronger alternative ASS
429	54bp upstream of exon 15 ASS. It is possible that this secondary competitor
430	further weakens the exon 15 ASS thus enhancing the effects of variant
431	chr8:g.87618576G>A.
432	Next, using OGI-081 as a true positive case, we tested whether splicing
433	prediction algorithms could be used to prioritize candidate non-coding splicing
434	altering variants, circumventing the need for RNA-seq analysis. We annotated

- the 3,268 rare variants with allelic pairs identified in OGI-081 with two splicing
- 436 prediction programs. (i). Alamut Batch that makes its prediction by the combined

437 calculations of the same splicing predictions algorithms as Alamut visual but is capable of calculating the effects of multiple variants. (ii). SpliceAl, a deep neural 438 network tool, to predict splice junctions from pre-mRNA transcript sequence ¹⁸. 439 Alamut batch calculated a high probability for altering splicing for 532 variants in 440 315 genes (Table S10). Although variant chr8:g.87618576G>A, the novel 441 442 pathogenic variant identified in this study, was predicted by Alamut Batch to strongly activate a cryptic donor site the large number of additional candidate 443 variants make this tool too cumbersome for identification of candidate non-coding 444 445 pathogenic variants. For SpliceAI, to identify synonymous exonic, near intronic, and deep intronic variants predicted to affect splicing at a validation rate of 40% 446 447 the authors used Δ Score greater than or equal to 0.2, 0.2, and 0.5 respectively. Out of the variants segregating in OGI-081 only eight had scores 0.2<0.5 and only 448 449 one, variant Chr9:g.86536129C>T, received a \triangle Score >0.5 (Table S11). Variant chr8:g.87618576G>A, the novel pathogenic variant identified in this study as 450 activating a cryptic donor splice site, was calculated by SpliceAI to have a donor 451 gain Δ Score of 0.3 well below the 0.5 cutoff for deep intronic variants. Thus, had 452 we used SpliceAI splicing predictions as a filter to identify potential causal 453 variants for functional validations, variant chr8:g.87618576G>A would have been 454 overlooked. The ASS of exon 14b was not identified by either algorithm and 455 456 therefore could not have been used to highlight variant chr8:g.87618576G>A as 457 a more plausible pathogenic variant.

458

459 **Discussion**

460 We present here an unbiased approach based on the combination of WGS and 461 RNA-seq data to identify and functionally validate pathogenic non-coding variants without the use of large datasets. We show that ex vivo models, such as iPSC 462 derived ROs, can serve as a surrogate source of a patient's own retinal tissue for 463 464 RNA and protein analyses. IRDs are currently at the focus of gene therapy advances and several clinical trials are underway, including a trial for CNGB3 465 gene augmentation therapy ⁶. This work was aimed at expanding the number of 466 patients eligible for clinical trials and forthcoming therapies. Indeed, our findings 467 here make the two affected siblings of OGI-081 eligible to participate in ongoing 468 clinical trials for CNGB3 gene therapy. Our approach is applicable to any 469 470 inherited disease, both WGS and RNA-seg techniques are commercially available, gold standards are being established and the analysis tools are readily 471 accessible ^{47,48,82,49–55,81}. Ex vivo organoid models are being developed for a 472 multitude of tissues including brain ^{83,84}, kidney ⁸⁵, liver ^{86,87} and lung ⁸⁸. 473 Non-coding variants present a challenge for a correct genetic diagnosis that is 474 imperative for a successful genetic therapy. The combination of WGS and RNA-475 476 seq methodologies allows us to both detect non-coding variants and evaluate their functionality throughout the genome. Indeed, a similar approach has already 477 been successfully employed to diagnose diseases where RNA could be 478 harvested from biopsies of disease-relevant tissue ^{19,20}. These studies relied on 479 the availability of large control datasets of RNA-seq samples from unaffected 480

individuals and/or a large cohort of patients ^{19,20}. Our work shows that the correct 481 diagnosis of non-coding variants is possible without reliance on such resources. 482 WGS analysis of all five members of OGI-081 and segregation analysis of the 483 variants within the family narrowed down the search from tens of thousands of 484 variants to a few hundred with allelic pairs. We then used RNA-seg analysis 485 486 comparing two affected siblings to an unaffected one as an orthogonal approach to identify genes with altered splicing or expression in disease. Thus identifying 487 the deep intronic variant chr8:g.87618576G>A as a novel pathogenic variant in 488 489 the CNGB3 gene. The iPSC derived ROs served both as a source of disease relevant transcriptome and as a system for functional validation of the truncated 490 proteins. In future studies, for families with a single patient the parents may serve 491 as control samples, so that each parent controls for the effect of the allele 492 inherited from the other parent making our approach applicable even for ultra-493 494 rare diseases.

Once the deep intronic variant was detected and validated we were able to use 495 that prior knowledge to identify additional factors that may have contributed to the 496 inclusion of exon 14b such as the availability of the cryptic acceptor site of exon 497 498 14b and the comparative weakness of the exon 15 acceptor site. Such complex 499 dependencies are a prime example as to why sequence based predictions of 500 splicing patterns are hard to compute. Still, several splicing predictors in the 501 Alamut Visual software were able to detect the increase in the splicing probability of the cryptic donor site as a result of variant chr8:g.87618576G>A. This 502 prompted us to test whether such splicing predictors can be used as preliminary 503

504 filters to identify candidate pathogenic variants for "gene by gene" validation methods, circumventing the need for RNA-seg analysis of ROs. We found that 505 the more established approach represented by the Alamut Batch method of 506 combining the calculations of several splicing predictors that are designed to 507 identify known splicing motifs yielded too many candidates for gene by gene 508 509 validation. In contrast, the more recent approach of deep neuronal networks algorithms, represented by SpliceAI, failed to assign high probability to the true 510 511 positive variant chr8:g.87618576G>A. Still, in cases where some prior knowledge 512 can help prioritize variants or highlight ones with lower than expected scores these methods may yet be helpful. In cases where no prior knowledge can help 513 prioritize candidate variants, such as in patients where both pathogenic alleles 514 are non-coding, and especially in cases where a cell type relevant for functional 515 validation is not available, the approach established here is preferable. 516

517 Appendices

518 Differential gene expression analysis

As mentioned above, differential gene expression analysis was less informative in the OGI-081 datasets. We compared gene expression levels from each of the affected siblings to that of the unaffected sibling (Table S9). We found 401 genes to be consistently down regulated, of which 27 were Y linked as expected given that the two affected siblings are females while the unaffected sibling is a male. We excluded these Y linked genes from further analysis. Tools are not currently available to filter out non-Y linked genes that may be differentially expressed

526 between the sexes under normal conditions in ROs. Of the remaining 374 down

- regulated genes, 29 also contained allelic variant pairs (Table S3). In addition,
- we found 1120 genes to be consistently up regulated between the two affected
- siblings and the unaffected sibling (Table S9). Of the up regulated genes, 15 also
- 530 contained allelic variant pairs (Table S3). None of the 44 differentially expressed
- 531 genes with allelic variant pairs are reported in RetNet as IRD genes.

532 Supplemental Data

533 Supplemental data includes two figures and 11 tables.

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- 550 ENCSR991HIR, ENCSR460YCS and ENCSR321PGV.

551 **Declaration of Interests**

552 The authors declare no competing interests.

553 Web Resources

- 554 ENCODE https://www.encodeproject.org/experiments/ENCSR551NII/
- 555 ExAC <u>http://exac.broadinstitute.org/</u>
- 556 gnomAD <u>http://gnomad.broadinstitute.org/</u>
- 557 GTEx Portal https://gtexportal.org/home/
- 558 Picard tools <u>http://broadinstitute.github.io/picard/</u>
- 559 RetNet <u>http://www.sph.uth.tmc.edu/RetNet/</u>

560 Accession Numbers

- 561 The accession numbers for the RNAseq samples reported in this paper
- 562 (BioProject: PRJNA564377) are:
- 563 SRA:SRR10082823
- 564 SRA:SRR10082822
- 565 SRA:SRR10082821

- 566 SRA:SRR10082828
- 567 SRA:SRR10082829
- 568 SRA:SRR10082824
- 569 SRA:SRR10082830
- 570 SRA:SRR10082827
- 571 SRA:SRR10082826
- 572 SRA:SRR10082825
- 573 SRA:SRR10082820

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866 Figure Legends

Figure 1: Family OGI-081 variant segregation scheme and retinal

- 868 **phenotypes.** A. The OGI-081 pedigree with variant segregation scheme. B.
- Fundus (upper panel) and OCT image (lower panel) for the OGI-081-197 at age
- 870 8, area of retinal degeneration is indicated by the red bar.
- Figure 2: RO differentiation. A. Schema of the differentiation process and a
- light microscopy image of a typical RO. Arrow head indicating photoreceptors,
- scale bar = 100 microns. B-G. Immunocytochemistry on cryosections of ROs.
- B&E. NR2E3 staining of rod nuclei (Green). C&F. Mature cones show staining of
- cone opsins in the cone photoreceptor outer segments (Red). C. S opsin. F. ML
- opsin. D&G. Overlay of rod and cone staining. All cones are stained with ARR3
- 877 (Purple). Scale bars = 20 microns.

878 **Figure 3: Comparison of IRD gene expression and splice junctions.** Human

- Normal Retina (HNR; N=3, gray), RO from the unaffected sibling (N=5, blue),
- Skin-Sun Exposed (SSE; N=473, green) and Whole Blood (WB; N=407, red).
- A&B Average TPM values of IRD genes. A. IRD genes are sorted by their
- expression in HNR overlaid with RO, SSE or WB. B. Violin plot. C&D Number of
- splice junctions detected by MAJIQ. C. All annotated genes. D. IRD genes.

Figure 4: Alternative splicing in the NCALD and CNGB3 genes. A. Venn

- diagram of genes found to have alternative splicing events in OGI-081
- comparison of affected vs. unaffected siblings and genes found to have
- segregating allelic pairs (green). Alternative splicing analysis was conducted by

MAJIQ (blue) and CASH (red). B. Collapsed diagram of exons (black boxes) from
all isoforms of the *NCALD* and *CNGB3* genes. DNA variants (red); MAJIQ (blue)
and CASH (yellow) alternative splicing events (E). Events detected by MAJIQ are
depicted as split reads arches. The event range detected by CASH is depicted by
the left (L) and right (R) borders. Genomic locations of variants, junctions and
event borders are given in table S6.

894 Figure 5: Aberrant splicing of CNGB3 in the affected vs. unaffected

siblings. A. Sashimi plot presenting RNA-seq results showing a cryptic exon 895 spliced into the isoform as a result of the intronic variant chr8:g.87618576G>A. 896 897 The cryptic exon is only present in the affected sibling (lower panel, red) and not in the unaffected sibling (upper panel, blue). The splice junction between exon 898 14b and exon 15 is not represented by split reads in the Sashimi plot due to an 899 900 alignment error (FigureS1). B. RT-PCR using primers from the canonical exon14 and exon 15. All three siblings express the normal size isoform lacking exon 14b 901 (lower band). A larger abnormal band containing exon14b (upper band) is 902 present in the two affected siblings Af1 and Af2 but not in the unaffected sibling 903 (Un). Negative controls lacking RNA template in the RT reaction (NC1) and NC1 904 used as template for PCR amplification (NC2). Sanger sequencing of the larger 905 906 band confirming the inclusion of exon 14b. C. Schematic representation of the protein domains in W.T CNGB3 and the two mutant alleles found in the affected 907 siblings of OGI-081. 908

Figure 6: Mislocalization of the CNGB3 truncated proteins.

910 Immunocytochemical analysis of day 262 ROs from the heterozygous parent

- 911 OGI-081-200 (A-C) and an affected sibling OGI-081-197 (D-F). In the
- 912 heterozygote, both ML opsins (red) and CNGB3 (green) are localized to the
- 913 photoreceptor outer segments whereas in the affected sibling, CNGB3 localizes
- to the photoreceptor inner segments. An exemplary photoreceptor outer segment
- is indicated by the white brackets. Nuclei are counterstained with DAPI (blue).
- 916 Scale bars = 20 micron

918 Table 1: Sample level alignment report and QC summary. A high average

SampleName	RawReads ^a	HQReads [♭]	% UniqAligned ^c
OGI-081-197-1	159,976,749	129,671,633	87.73
OGI-081-197-2	256,399,765	234,701,223	90.52
OGI-081-197-3	166,237,226	155,442,261	88.88
OGI-081-198-1	167,327,080	145,058,230	88.66
OGI-081-198-2	197,037,935	179,026,100	89.13
OGI-081-198-3	191,485,904	177,663,416	90.15
OGI-081-340-1	175,199,604	156,566,678	91.01
OGI-081-340-2	227,928,431	216,526,830	89.53
OGI-081-340-3	137,096,260	129,192,437	90.47
OGI-081-340-4	151,572,341	140,982,711	90.59
OGI-081-340-5	164,908,758	154,550,198	99.66

919 unique percentage alignment rate is reported.

920

^a Raw reads count (RawReads), ^b Filtered high quality reads (HQReads), ^c

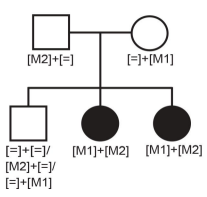
921 Percent of uniquely aligned reads (% UniqAligned).

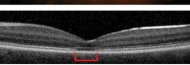
Table 2: Splicing junctions involved with the inclusion of exon14b.

	Exon 14 Donor site	Exon 14b Acceptor site	Exon 14b Donor site		Exon 15 Acceptor site	
	Annotated	W.T	W.T	Variant	Annotated	W.T
NM_019098.4 ^a	c.1662	c.1663-2238	c.1663- 2137	c. 1663- 2137C>T	c.1663	c.1663- 54
SSF⁵	95.3	83.3	67.8	73.7	76.7	90.4
MaxEntScan ^c	10.5	4.8	0	4.8	6.1	8.7
NNSplice ^d	1	0.5	0	1	0.1	0.8

- ^a The NM_019098.4 isoform of CNGB3 is used to define the cDNA coordinates. ^{b-}
- ^d Splicing prediction scores are given from the three algorithms, SSF,
- 925 MaxEntScan and NNSplice.

Α.

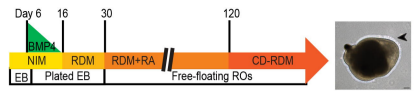


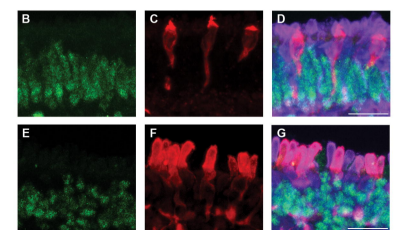


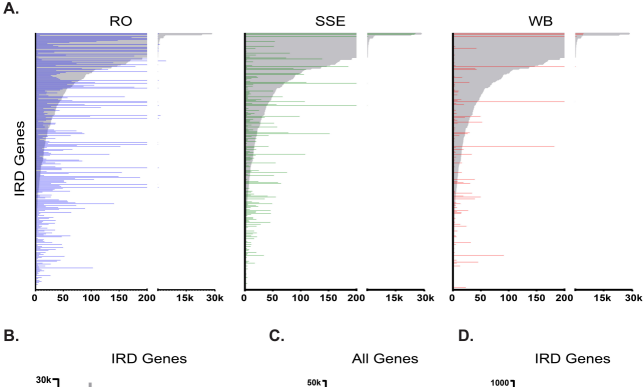


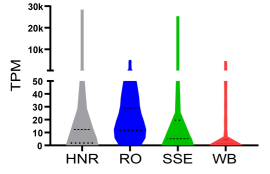
Β.

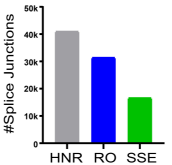
Α.

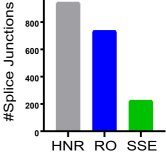












MAJIQ CASH 13 233 82 NCALD & CNGB3 12 4 624 Allelic pairs genes

Α.

