

# AAV-mediated gene augmentation therapy of *CRB1* patient-derived retinal organoids restores the histological and transcriptional retinal phenotype

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## SUMMARY

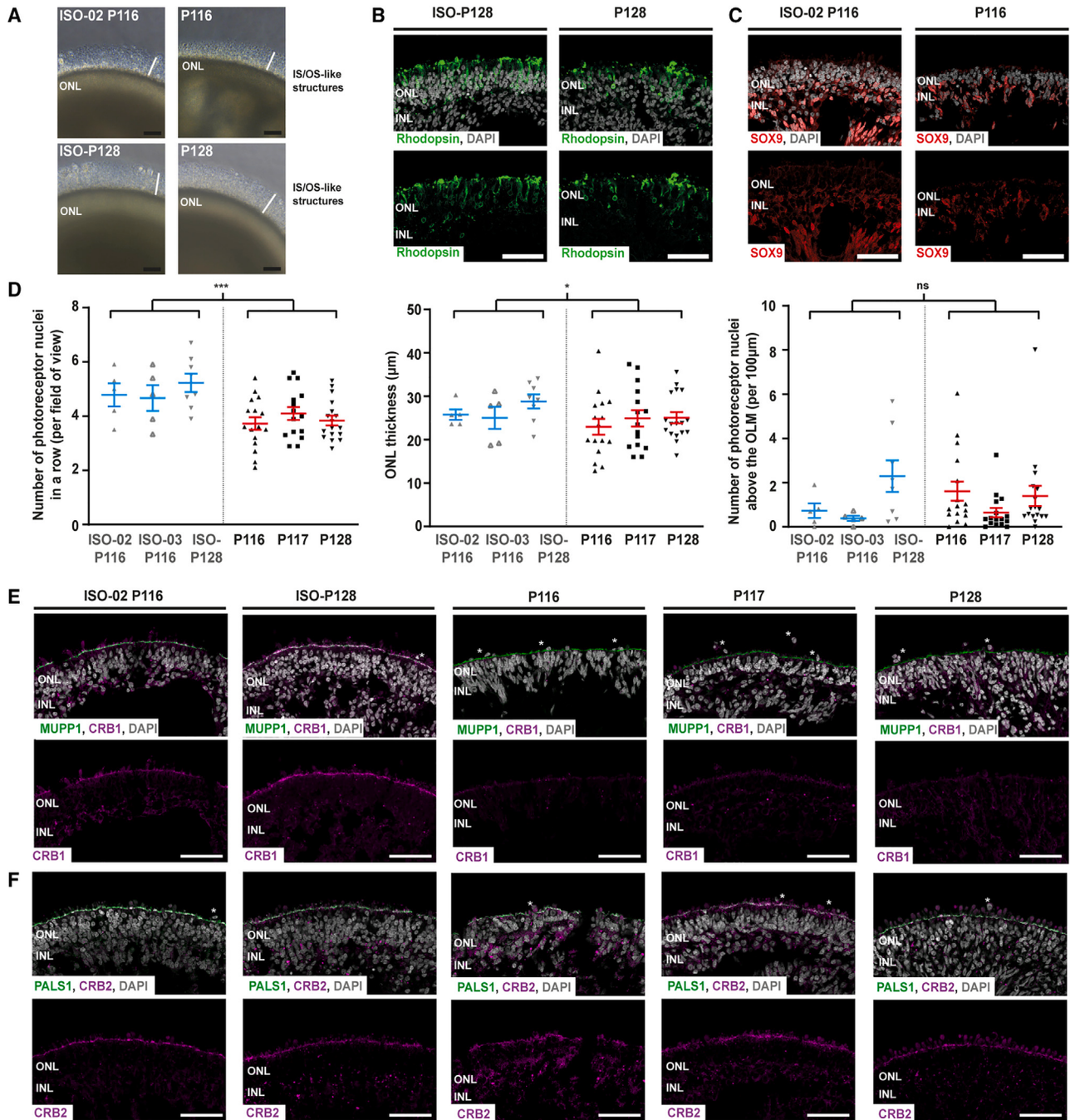
Retinitis pigmentosa and Leber congenital amaurosis are inherited retinal dystrophies that can be caused by mutations in the Crumbs homolog 1 (*CRB1*) gene. *CRB1* is required for organizing apical-basal polarity and adhesion between photoreceptors and Müller glial cells. *CRB1* patient-derived induced pluripotent stem cells were differentiated into *CRB1* retinal organoids that showed diminished expression of variant *CRB1* protein observed by immunohistochemical analysis. Single-cell RNA sequencing revealed impact on, among others, the endosomal pathway and cell adhesion and migration in *CRB1* patient-derived retinal organoids compared with isogenic controls. Adeno-associated viral (AAV) vector-mediated *hCRB2* or *hCRB1* gene augmentation in Müller glial and photoreceptor cells partially restored the histological phenotype and transcriptomic profile of *CRB1* patient-derived retinal organoids. Altogether, we show proof-of-concept that AAV.*hCRB1* or AAV.*hCRB2* treatment improved the phenotype of *CRB1* patient-derived retinal organoids, providing essential information for future gene therapy approaches for patients with mutations in the *CRB1* gene.

## INTRODUCTION

Retinitis pigmentosa (RP) and Leber congenital amaurosis (LCA) are inherited retinal dystrophies caused by mutations in, among others, the Crumbs homolog 1 (*CRB1*) gene (Den Hollander et al., 1999; Nguyen et al., 2022; Talib et al., 2017). Canonical *CRB1* is a large transmembrane protein consisting of a short 37 amino acid intracellular domain containing a protein 4.1, ezrin, radixin, moesin (FERM), and a conserved glutamic acid-arginine-leucine-isoleucine (ERLI) PDZ binding motif, a single transmembrane domain, a large extracellular domain with multiple epidermal growth factor and laminin-A globular-like domains (Boon et al., 2020; Den Hollander et al., 1999; Quinn et al., 2017). Recently, a short non-canonical alternatively spliced form of *CRB1*, *CRB1-B*, has been described containing substantial extracellular domain overlap but with distinct amino terminus and lacking the C-terminal transmembrane and intracellular domains (Ray et al., 2020). In mammals, canonical *CRB1* is a member of the Crumbs family together with *CRB2* and *CRB3A*. The canonical *CRB* complex is formed by interaction with protein associated with Lin seven 1 (PALS1), also known as membrane-associated guanylate kinase p55 subfamily member 5 (MPP5), which binds to the conserved C-terminal PDZ domain of *CRB* (Margolis, 2018; van de Pavert et al., 2004; Roh et al., 2002). Binding of PALS1 can lead to the recruitment of multiple PDZ domain protein 1 (MUPP1) or the InaD-

like protein (INADL/PAT) to the apical membrane (Roh et al., 2002). This *CRB* complex is evolutionary conserved and is important for maintaining cell adhesion and regulating apical-basal polarity (Bulgakova and Knust, 2009).

So far, no treatment possibilities are available for patients with RP or LCA caused by mutations in *CRB1*. Gene augmentation therapies using adeno-associated viral (AAV) vectors are of emerging interest for retinal dystrophies because of the recent FDA approval of an AAV-mediated gene therapy approach for RP and LCA patients with mutations in the *RPE65* gene (Maguire et al., 2019). AAVs are the leading platform for gene delivery because of their low toxicity, limited integration into the host genome, and because different AAV capsids display distinct cell tropisms. Their major disadvantage is the limited packaging capacity; inverted terminal repeats, cDNAs, and regulatory sequences bigger than 4.9 kb often do not fit in a single AAV capsid. Unfortunately, the full-length cytomegalovirus (CMV) ubiquitous promoter and *hCRB1* cDNA exceeds this packaging limit. However, substantial expression levels of canonical *hCRB1* protein in mouse mutant *Crbl* retina were observed using an AAV with codon optimized *hCRB1* cDNA linked to a minimal CMV promoter (Pellissier et al., 2014a). This AAV.CMVmin.*hCRB1* was deleterious upon intravitreal injection in *Crbl* mouse models (Pellissier et al., 2015). As an alternative approach, *CRB* family member *CRB2* was used to restore retinal function and vision in *Crbl* mice (Buck et al., 2021; Pellissier et al., 2015), showing



### Figure 1. *CRB1* patient and isogenic control phenotypic analysis at DD210

(A) Representative bright-field images of ISO-02 P116, P116, ISO-P128, and P128 cultured organoids.

(B) Representative immunohistochemical images of rhodopsin (green) in ISO-P128 and P128.

(C) Representative immunohistochemical images of SOX9 (red) in ISO-P116 and P116.

(D) Quantitative analysis of number of photoreceptor nuclei in a row per field of view ( $p = 0.000$ ), ONL thickness per field of view ( $p = 0.049$ ), and number of photoreceptor nuclei above the OLM per  $100\mu\text{m}$  ( $p = 0.651$ ) in *CRB1* patient-derived and isogenic control retinal organoids.

(E) Immunohistochemical images of CRB1 (red) and MUPP1 (green) in *CRB1* patient-derived retinal organoids with two appropriate isogenic controls.

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the potential of AAV.h*CRB2* gene augmentation therapy for patients with mutations in the *CRB1* gene.

There are several mouse models described with mutations in the *Crb1* and/or *Crb2* gene mimicking the RP or LCA phenotype (Alves et al., 2013, 2014, 2019; Mehalow et al., 2003; van de Pavert et al., 2004, 2007a; Pellissier et al., 2013; Quinn et al., 2018, 2019a). However, immunoelectron microscopy identified the subcellular localization of CRB1 and CRB2 proteins to be different in mouse and human models. In mice, CRB2 is present in photoreceptor cells and Müller glial cells (MGCs) at the subapical region (SAR) of the outer limiting membrane (OLM), while CRB1 is solely present in MGCs at the SAR (van Rossum et al., 2006). In contrast, in human fetal retina and human induced pluripotent stem cell (hiPSC)-derived retinal organoids both CRB1 and CRB2 are observed at the SAR in photoreceptors and MGCs (Quinn et al., 2019b). This discrepancy suggests the importance of using human-derived models for gene therapy approaches.

Here, we describe the phenotype of differentiation day 210 (DD210) and DD230 patient-derived retinal organoids harboring *CRB1* missense mutations compared with isogenic controls in more detail using immunohistochemical analysis and single-cell RNA sequencing (scRNA-seq). Next, the effect of AAV-mediated h*CRB1* or h*CRB2* gene augmentation therapy was analyzed on *CRB1* patient-derived and isogenic control retinal organoids. A partially improved retinal phenotype of *CRB1* patient-derived retinal organoids was observed, providing crucial data for future gene therapy approaches for patients with mutations in the *CRB1* gene.

## RESULTS

### Reduced number of photoreceptor nuclei and thinner outer nuclear layer in DD210 *CRB1* patient-derived retinal organoids compared with isogenic controls

Retinal organoids were differentiated from hiPSC lines derived from three *CRB1* RP patients: (1) LUMC0116iCRB with c.3122T>C p.(Met1041Thr) homozygote missense mutations (here abbreviated as P116), (2) LUMC0117iCRB with 2983G>T p.(Glu995\*) and c.1892A>G, p.(Tyr631Cys) mutations (P117), and (3) LUMC0128iCRB with c.2843G>A p.(Cys948Tyr) and c.3122T>C p.(Met1041Thr) missense mutations (P128) (Quinn et al., 2019b). Isogenic controls

for P116 and P128 were generated by CRISPR-Cas9; (1) ISO-02 P116 with a homozygous correction, (2) ISO-03 P116 with a heterozygous correction, and (3) ISO-P128 a heterozygous correction of Cys948Tyr (Table S1). Genomic stability of all iPSC lines was tested by a digital PCR test of the copy number variants (CNVs) of 90% of the most recurrent abnormalities in hiPSCs (Assou et al., 2020). No aberrant CNVs were observed in the hiPSC lines used in this study (Figure S1A).

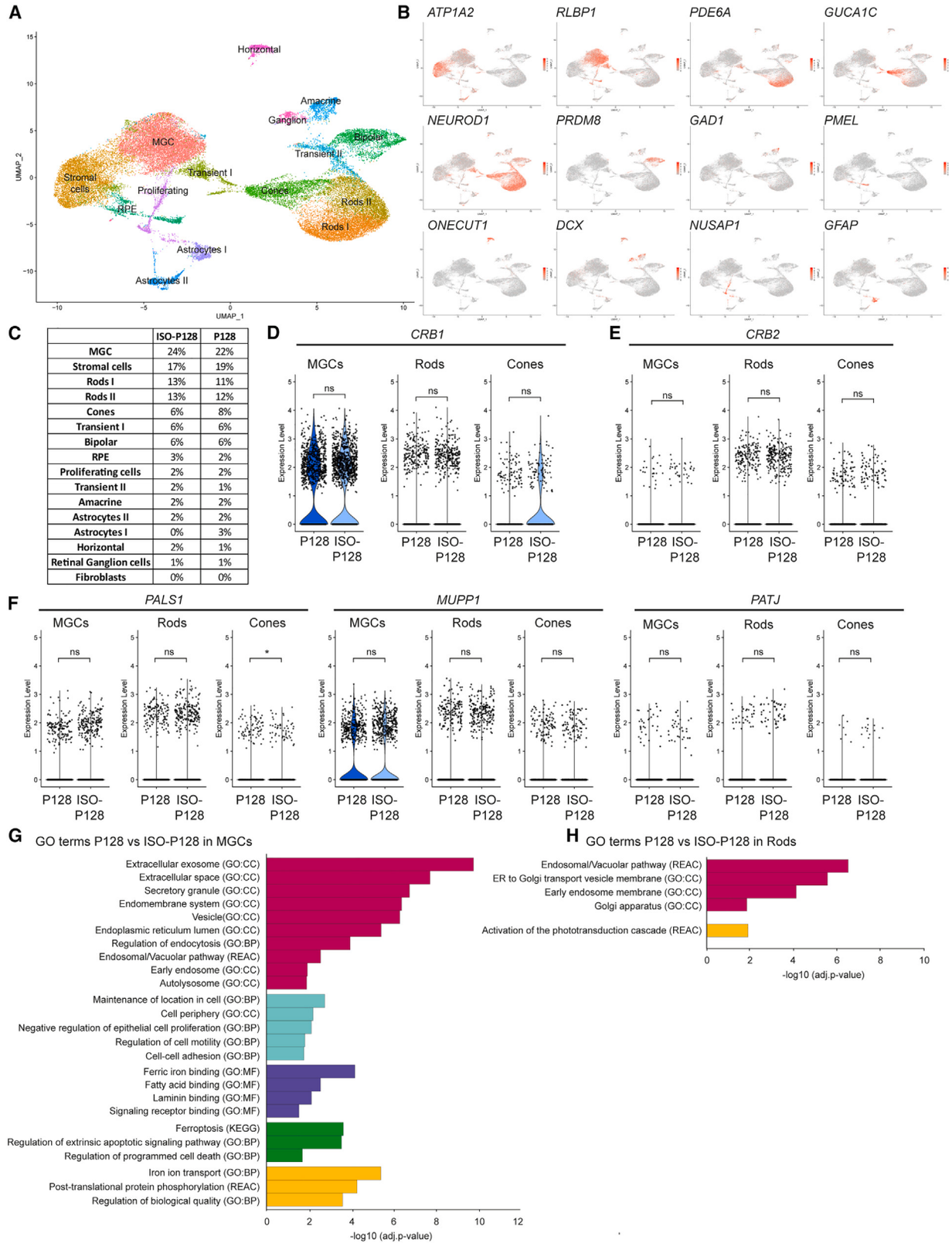
Previous research has shown that *CRB1* patient-derived retinal organoids at DD180 show disruptions at the OLM and photoreceptor nuclei protruding above the OLM (Quinn et al., 2019b). Here, we analyzed the phenotype of the *CRB1* patient-derived retinal organoids at a later timepoint (DD210) and compared those with the isogenic controls. By light microscopy, no visible difference was observed in cultured retinal organoids comparing *CRB1* patient with the isogenic controls at DD210: all contained a translucent region at the outside of the organoid (the ONL) with inner and/or outer-segment-like structures around the retinal organoid (Figures 1A and S1B). Immunohistochemistry of rod photoreceptor marker rhodopsin and MGC marker SOX9 at DD210 revealed the presence of rod photoreceptors and SOX9-positive MGCs for both patient and isogenic controls (Figures 1B, 1C, S1C, and S1D).

When analyzing the phenotype in more detail, a moderate but statistically significant decrease in the number of photoreceptor nuclei and ONL thickness was observed in *CRB1* patient-derived retinal organoids compared with isogenic controls (Figure 1D). In contrast to *CRB1* patient-derived retinal organoids at DD180 (Quinn et al., 2019b), no statistically significant difference in the number of photoreceptor nuclei above the OLM was detected at DD210 (Figure 1D). In addition, no statistically significant differences were observed for the total retinal thickness and the inner nuclear layer (INL) thickness of the retinal organoids (Figure S1H).

### Missense mutations in *CRB1* result in reduced levels of variant *CRB1* protein in *CRB1* patient-derived retinal organoids

Immunohistochemistry analysis of all three *CRB1* patient-derived retinal organoids at DD210 shows a diminished *CRB1* staining at the OLM compared with the isogenic

(F) Immunohistochemical images of CRB2 (red) and PALS1 (green) in *CRB1* patient-derived retinal organoids with two appropriate isogenic controls. Each datapoint in the graph represents individual organoids, of which an average has been taken of 3–6 representative images. The standard error of mean (SEM) is derived from these averages. Number of individual organoids per condition and differentiation round: P116, n = 16; P117, n = 15; P128, n = 17, from four independent organoid batches; ISO-P128, n = 8 from three independent organoid batches; ISO-02 P116, n = 5; and ISO-03 P116, n = 5 from two independent organoid batches. Scale bars, 50  $\mu$ m. Statistical analysis: generalized linear mixed models with \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001. Related to Figure S1.



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control (Figures 1E and S1E). Similar strongly diminished levels of variant *CRB1* in *CRB1* patient-derived retinal organoids compared with the isogenic control were observed at DD180 (Figure S1G), while *CRB2* and *CRB* complex members *MUPP1* and *PALS1* remain at the OLM in both isogenic and *CRB1* patient-derived organoids (Figures 1E, 1F, and S1F).

### Missense mutations in *CRB1* do not affect the levels of *CRB1* or *CRB2* RNA transcripts in *CRB1* patient-derived retinal organoids

Next, we used scRNA-seq to identify differences in RNA transcripts and Gene Ontology (GO) pathways between DD230 *CRB1* patient-derived retinal organoids and isogenic controls. Transcriptionally similar cells were grouped and visualized (R package Seurat), revealing distinct clusters with differentially expressed marker genes (Table S3). These identified expressed genes per cluster were compared with known retinal marker genes to classify clusters. Major retinal cell types could be visualized on a UMAP plot, such as MGCs, photoreceptor cells (both rods and cones), bipolar cells, amacrine cells, horizontal cells, ganglion cells, and retinal pigment epithelium (RPE) (Figure 2A). In addition, some of the clusters consisted of astrocytes and transitory cells (Figure 2A), and tissue that is generally attached to the retinal organoid was observed and classified as stromal cells (Figures 2A and S2C). The expression of key cell-type-specific markers of all clusters are shown in a feature plot and heatmap (Figures 2B and S2A). Interestingly, two rod photoreceptor cell subtypes can be distinguished. Upon further analysis, cluster rods I was identified to be composed of more mature cells with significantly higher transcript expression levels of *NR2E3*, *PDE6B*, and *RHO* in comparison with cluster rods II (Figure S2B) (Swaroop et al., 2010).

We confirmed that all the major retinal cell clusters were equally present in both P128 and ISO-P128 (Figure 2C) and P116 and ISO-P116 (Figure S3A). Expression levels of both *CRB1* and *CRB2* were predominantly observed in MGCs and photoreceptor cells. When analyzing *CRB1* expression levels in MGCs and photoreceptor cells in more detail, no

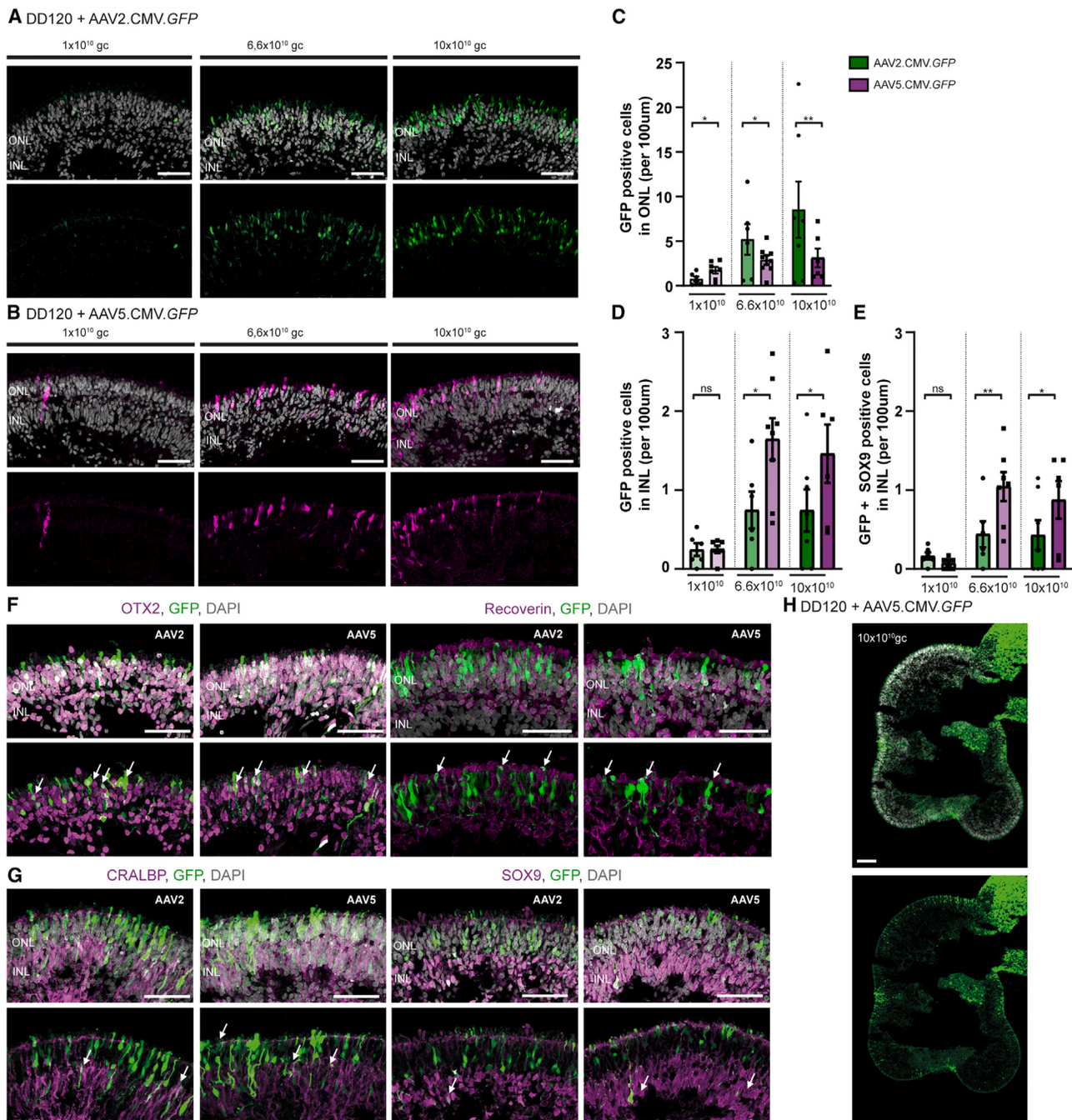
statistically significant differences were observed between P128 and ISO-P128 (Figure 2D) or between P116 and ISO-P116 (Figure S3B). Rods I and rods II were combined into a general “rods” cluster, since no statistically significant differences were observed in the individual clusters (data not shown). Moreover, the sequence of *CRB1-B*, an alternative transcript of *CRB1* containing distinct 5' and 3' ends, was added to the pre-built reference and was not detected in our DD230 retinal organoids (Figure S2D). While *CRB1-B* transcripts were detected in adult human retina cDNA, the levels of *CRB1-B* transcripts were below detection level by quantitative reverse transcription real-time PCR (qRT-PCR) on DD210 retinal organoids (data not shown; experimental procedures in supplemental information). In addition, no statistically significant difference was observed for the *CRB2* expression level (Figures 2E and S3C), canonical *CRB* core complex members *PALS1*, *MUPP1*, *PATJ* (Figures 2F and S3D), or FERM proteins *MSN*, *EZR*, or *EPBA1L5* (Figure S2E).

### Gene profiling shows disruptions in the endosomal system in MGCs and rods

Differential gene expression analysis followed by GO term analysis comparing *CRB1* patient and isogenic control retinal organoids was performed. Analysis of P128 and ISO-P128 specifically in the MGCs, where most *CRB1* is expressed, revealed five groups of similar GO terms containing differentially expressed genes deregulated in the patient-derived retinal organoids. The first GO group is involved in the endosomal system, including extracellular exosomes, vesicles, endomembrane system, and early endosomes (Figure 2G). The second group is involved in the maintenance of location in the cell, cell motility, proliferation, and cell-cell adhesion, the third group revealed differences in proteins containing various binding domains such as ferric iron and fatty acid, while the fourth group is involved in cell death (Figure 2G). Finally, the last one is a mixed group with pathways such as iron ion transport and post-translational protein phosphorylation (Figure 2G). In addition, as the *CRB1* transcript is also present in photoreceptor cells, differentially expressed markers

## Figure 2. scRNA-seq analysis comparing ISO-P128 with P128 shows disruptions in the endosomal system in Müller glial cells and rods

- (A and B) (A) UMAP plot of observed clusters and (B) expression plots of top markers indicating the distinct clusters.  
(C) Table showing that all retinal cell types are present in both lines.  
(D) Violin plots of *CRB1* transcript levels specifically in MGCs ( $p = 0.93$ ), rods ( $p = 0.84$ ), and cones ( $p = 0.72$ ).  
(E) Violin plots of *CRB2* transcript levels specifically in MGCs ( $p = 0.43$ ), rods ( $p = 0.18$ ), and cones ( $p = 0.85$ ).  
(F) Violin plots of canonical core *CRB* complex members *PALS1*, *MUPP1*, and *PATJ* transcript levels in MGCs ( $p = 0.32$ ;  $p = 0.51$ ;  $p = 0.73$ , respectively), rods ( $p = 0.97$ ;  $p = 0.18$ ;  $p = 0.63$ ), and cones ( $p = 0.025$ ;  $p = 0.25$ ;  $p = 0.87$ ).  
(G and H) Gene ontology (GO) analysis of differentially expressed markers specifically in MGCs (G) and rods (H) clustered in groups with similar terms in the same color. Number of independent organoids used: ISO-P128,  $n = 6$ ; and P128,  $n = 6$ , from one differentiation round equally divided into three separate sequencing rounds. Related to Figures S2 and S3 and Table S4.



**Figure 3. AAV2.CMV.GFP and AAV5.CMV.GFP transduction of DD120 control retinal organoids**

(A and B) Representative immunohistochemical images of (A) AAV2.CMV.GFP- and (B) AAV5.CMV.GFP-treated control organoids at DD120 with three different titer concentrations:  $1 \times 10^{10}$ ,  $6.6 \times 10^{10}$ , and  $10 \times 10^{10}$  genome copies (gc).

(C–E) Quantification of AAV-treated retinal organoids with AAV2.CMV.GFP or AAV5.CMV.GFP at the (C) ONL, (D) INL, or (E) GFP-positive MGC in the INL.

(F and G) Representative immunohistochemical images of photoreceptor cell markers (OTX2 and recoverin) and MGC markers (CRALBP and SOX9) showing colocalization with AAV.GFP for both AAV2.CMV.GFP- and AAV5.CMV.GFP-treated organoids.

(H) Representative 10× magnification immunohistochemical analysis of DD120 control organoid transduced with  $10 \times 10^{10}$  gc AAV5.CMV.GFP. Immunohistochemical images of (F, G, and H) are merged z stack views, the others are single image views. Scale bars, 50 µm. Each datapoint in the graph represents individual organoids, of which an average has been taken of at least three representative

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and subsequent GO terms were analyzed in rods (combination of rods I and rods II) and cones. In rods, GO terms involved in the endosomal system were observed to be differentially expressed (Figure 2H). In addition, markers associated with the activation of the phototransduction cascade were detected (Figure 2H). No statistically significant differentially expressed markers were observed in cone photoreceptor cells (data not shown).

Such an analysis was also performed comparing P116 with ISO-P116, where GO terms associated with the endosomal system were differentially expressed in rod photoreceptor cells (Figure S3E). No statistically significant GO terms were observed in MGCs, explained by the low number of differentially expressed genes and the low number of cells sequenced in this cluster. Altogether, these data show aberrations in the endosomal system between *CRB1* patient-derived retinal organoids compared with their isogenic controls.

#### Serotype AAV5.CMV.GFP is more efficient than AAV2.CMV.GFP in transducing MGCs at DD120

*CRB1* protein is localized at the OLM in human and non-human primate MGCs and photoreceptors (Quinn et al., 2019a, 2019b), and higher levels of *CRB1* transcript are found in MGCs than in photoreceptors (Figure 2D). For AAV-mediated gene therapy approaches in *CRB1* patient-derived retinal organoids it is therefore essential to transduce a sufficient number of MGCs in addition to photoreceptors. Therefore, we identified which cells are transduced using specific viral capsids and titers at DD120. Control retinal organoids were transduced with  $1 \times 10^{10}$  genome copies (gc),  $6.6 \times 10^{10}$  gc, or  $10 \times 10^{10}$  gc AAV2/5.CMV.GFP (AAV5.CMV.GFP) or AAV2/2.CMV.GFP (AAV2.CMV.GFP) and analyzed using immunohistochemistry after 3 weeks in culture.

A significant dose-dependent increase of GFP-positive cells was observed when control organoids were treated with AAV5.CMV.GFP or AAV2.CMV.GFP at DD120 (Figures S3F and S3G). The AAV-treated retinal organoids were quantified for number of GFP-positive cells in the ONL, the INL, and GFP-positive cells in the INL that were also SOX9 positive (marking MGCs). AAV2.CMV.GFP transduced more photoreceptor cells in the ONL than AAV5.CMV.GFP (Figure 3C). However, AAV5.CMV.GFP transduced more cells in the INL than AAV2.CMV.GFP (Figure 3D). More specifically, more SOX9-positive MGCs were transduced with AAV5.CMV.GFP than with AAV2.CMV.

GFP (Figure 3E). Co-staining with photoreceptor markers (OTX2 and recoverin) and MGC markers (CRALBP and SOX9) confirmed the transduction of both cell types in AAV2.CMV.GFP as well as AAV5.CMV.GFP transduced organoids at DD120 (Figures 3F and 3G). Moreover, a  $10\times$  magnification of a retinal organoid treated with  $10 \times 10^{10}$  gc AAV5.CMV.GFP at DD120 showed that most of the retinal organoid was transduced in our experiment (Figure 3H).

Because AAV5 transduced more MGCs than AAV2, treatment with AAV5 at DD120 was used for further AAV.hCRB gene augmentation therapy with an intermediate dose of  $3.3 \times 10^{10}$  gc.

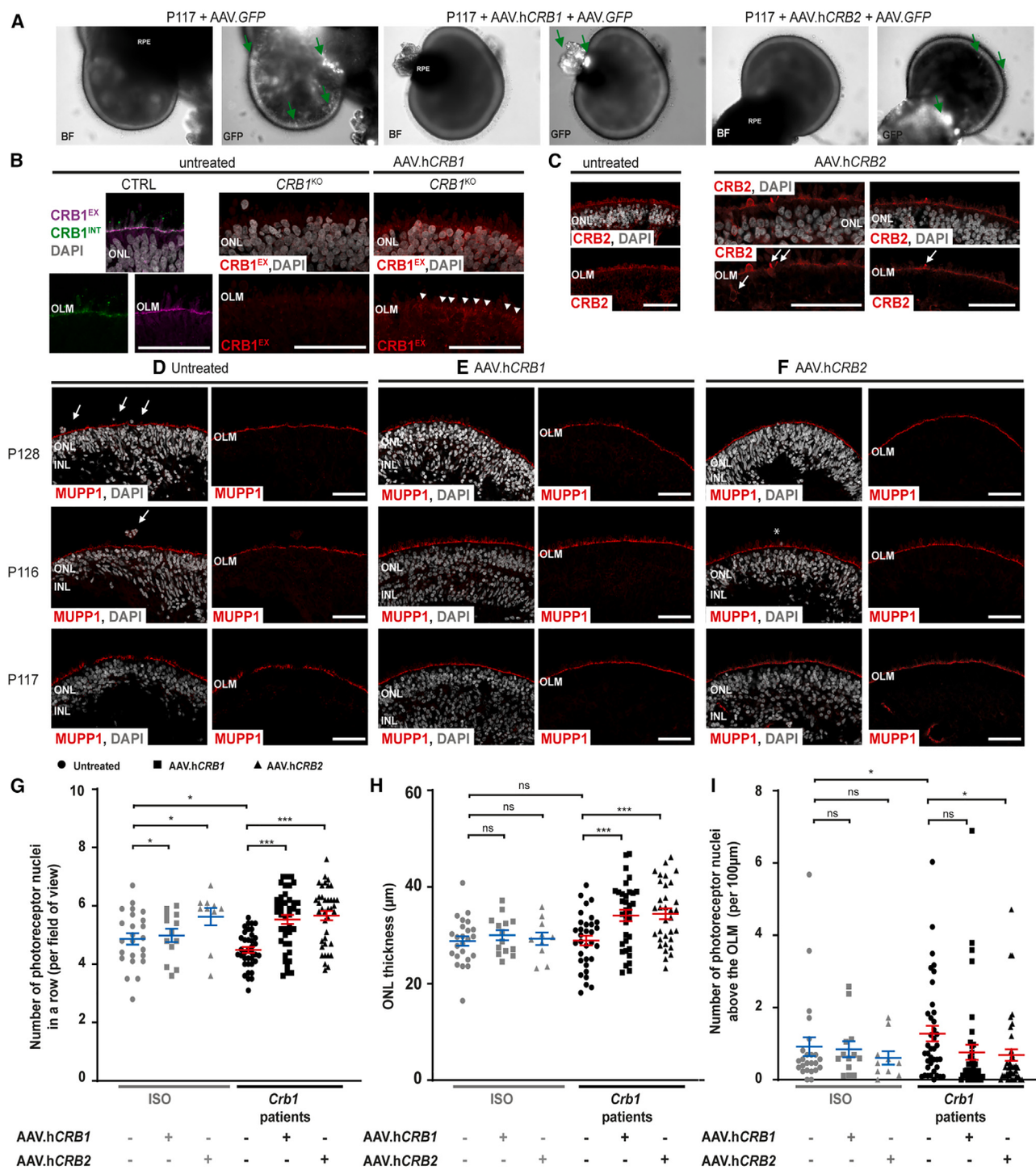
#### AAV-mediated hCRB gene augmentation therapy partially restores the histological phenotype of *CRB1* patient-derived retinal organoids

After defining the AAV.GFP tropism, preclinical gene therapy approaches were performed on *CRB1* patient-derived retinal organoids using AAV5.CMVmin.hCRB1 or AAV5.CMV.hCRB2 (here abbreviated as AAV.hCRB1 and AAV.hCRB2, respectively) at DD120 and analyzed at DD180 or DD210.

Immunohistochemical analysis and subsequent quantification of retinal organoids transduced with AAV.hCRB1 or AAV.hCRB2 and analyzed at DD180 show an increased number of photoreceptor nuclei in a row in AAV.CRB-treated P117 compared with the control (Figure S4B). In addition, fewer photoreceptor nuclei protruding above the OLM were observed after AAV.CRB treatment of P117 compared with the control organoids (Figure S4C). No statistically significant difference was observed for the retinal or the ONL thickness in both *CRB1* patient and control organoids treated with AAV.hCRB1 or AAV.hCRB2 (Figures S4D and S4E).

In addition, the long-term gene augmentation effect was examined for multiple *CRB1* patient-derived lines, where the organoids were collected and analyzed at DD210. One group of three different *CRB1* patient-derived retinal organoids was treated solely with AAV.hCRB1 or AAV.hCRB2 or left untreated, while in the following experiment the *CRB1* patient-derived retinal organoids were treated with AAV.hCRB1 with AAV.GFP, or AAV.hCRB2 with AAV.GFP, or AAV.GFP alone. Adding AAV.GFP facilitates in defining the regions where the AAV.hCRB most likely infected. Fluorescent images of organoids in culture co-treated with AAV.GFP and AAV.hCRB show the presence of GFP-positive regions, while no

images. The standard error of mean (SEM) is derived from these averages. Number of individual organoids per condition: for AAV2.CMV.GFP  $1 \times 10^{10}$  n = 5,  $6.6 \times 10^{10}$  n = 6, and  $10 \times 10^{10}$  n = 7, and for AAV5.CMV.GFP  $1 \times 10^{10}$  n = 7,  $6.6 \times 10^{10}$  n = 8, and  $10 \times 10^{10}$  n = 6 individual organoids from two independent differentiation rounds. Statistical analysis: generalized linear mixed models with \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001. Related to Figure S3.



**Figure 4. AAV-mediated gene therapy treatment on *CRB1* patient-derived and isogenic control retinal organoids**

(A) Representative bright-field (BF) and fluorescent (GFP regions indicated with green arrow) images of DD210 cultured P117 retinal organoids treated with  $3.3 \times 10^{10}$  vg AAV.hCRB.

(B) Immunohistochemical image of CRB1 in an untreated control retinal organoid, an untreated *CRB1*<sup>KO</sup> retinal organoid, and an AAV.hCRB1-treated *CRB1*<sup>KO</sup> retinal organoid showing increased CRB1 localization at the OLM of AAV.hCRB1-treated *CRB1*<sup>KO</sup> retinal organoids (arrowheads).

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visible differences were observed between treated and untreated organoids using bright-field or fluorescent images (Figure 4A). Immunohistochemical staining of CRB1 in AAV.hCRB1 or CRB2 in AAV.hCRB2-treated retinal organoids showed proof of recombinant CRB protein localization at the OLM and in the RPE (Figures 4B, 4C, and S4F–S4I). Further immunohistochemical analysis showed a partial improvement in the observed phenotype after AAV.hCRB1 or AAV.hCRB2 treatment (Figures 4D–4F). For quantitative analysis, all three *CRB1* patient-derived retinal organoids with and without concomitant treatment of AAV.GFP were pooled. No statistically significant difference was observed in fluorescence intensity of MUPP1 at the OLM in untreated compared with AAV.hCRB-treated *CRB1* patient-derived retinal organoids (Figure S4J). In addition, the expression of another core CRB-complex member, PALS1, is not changed after AAV.hCRB treatment (Figure S4K).

A statistically significant increased number of photoreceptor nuclei in a row was detected after AAV.hCRB1 and AAV.hCRB2 treatment at DD210, while this large difference was not observed in the treated isogenic controls (Figure 4G). Moreover, the ONL thickness (but not the retinal or the INL thickness) was significantly increased after AAV.hCRB treatment of *CRB1* patient-derived retinal organoids (Figures 4H, S4L, and S4M). Finally, the number of photoreceptor nuclei above the OLM was significantly improved after AAV.hCRB2 treatment of *CRB1* patient-derived retinal organoids at DD210 (Figure 4I). No statistically significant improvement in the number of photoreceptor nuclei above the OLM was observed after AAV.hCRB1 treatment of *CRB1* patient-derived retinal organoids nor after AAV.hCRB1 or AAV.hCRB2 treatment in the control retinal organoids (Figure 4I).

Altogether, these data show that the phenotype observed at DD180 and DD210 in *CRB1* patient-derived retinal organoids can be partially restored using AAV.hCRB1 or AAV.hCRB2 treatment.

### Differentially expressed genes related to the endosomal system are partially restored in AAV.hCRB-treated *CRB1* patient-derived retinal organoids

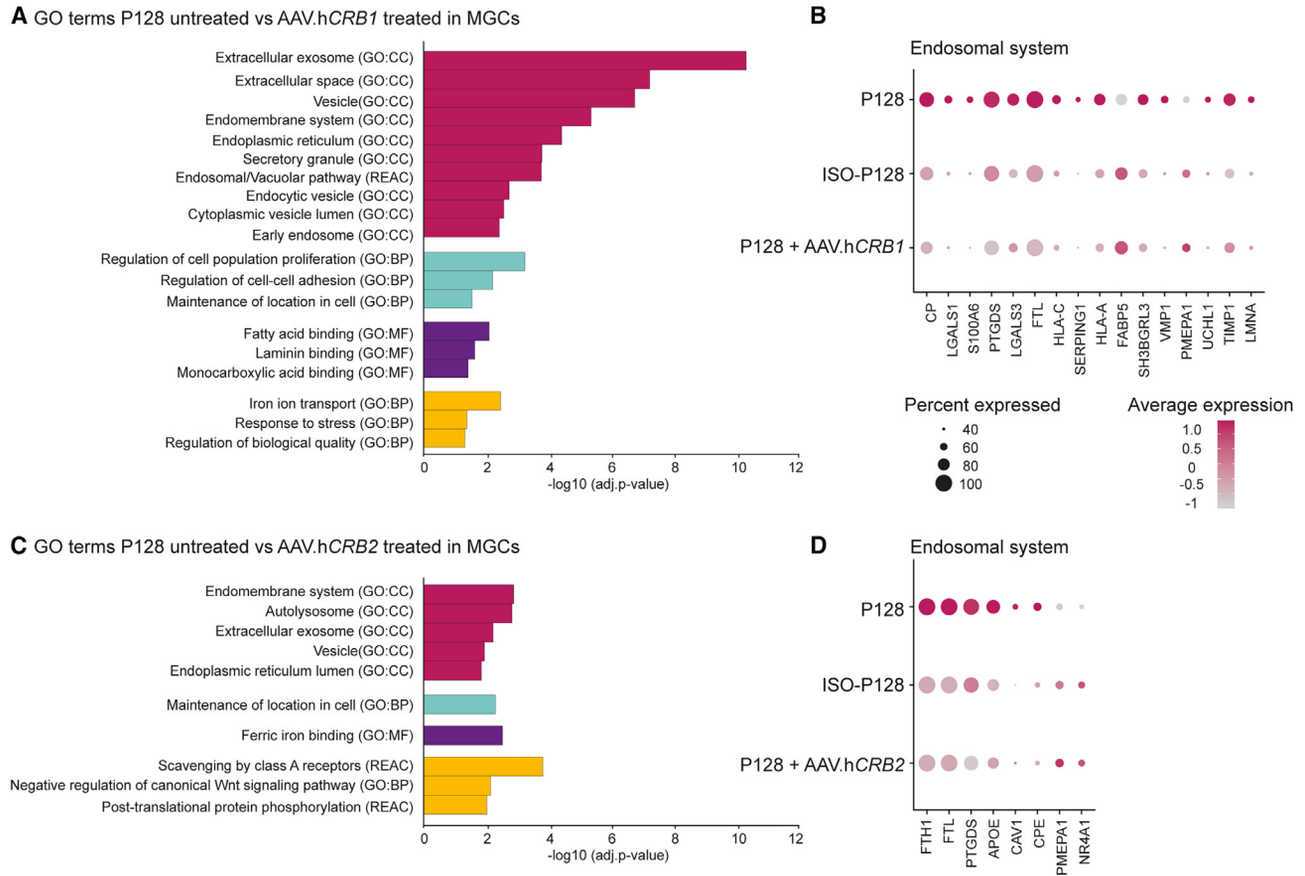
To identify gene expression changes upon *CRB* gene augmentation therapy, all three *CRB1* patient-derived retinal organoids were treated with AAV.hCRB1 and AAV.GFP or AAV.hCRB2 and AAV.GFP and compared with the AAV.GFP-treated control at DD230 using scRNA-seq. For all three patient-derived retinal organoids, we confirmed that the major retinal cell clusters were equally present in untreated and AAV-treated conditions (data not shown). A custom reference with the AAV.GFP, codon optimized AAV.hCRB1, and codon optimized AAV.hCRB2 sequences was added to the dataset to detect which cell clusters were transduced. While analyzing all organoids and conditions together, we observed that AAV.GFP mainly transduces RPE, photoreceptor cells, transient I, and MGCs (Figure S5A). Specifically, 66% of the RPE, 35% of rods, 36% of cones, 37% of transient I, and 20% of MGCs contained AAV.GFP expression. This is in line with what we observed previously in the immunohistochemical analysis (Figures 3E and 3F). Next, AAV.hCRB1 and AAV.hCRB2 expression was analyzed in AAV.hCRB-treated retinal organoids. While being unable to fully distinguish exogenous and endogenous hCRB due to the high sequence similarity and the low levels of endogenous *CRB1* and *CRB2* in DD230 RPE, we observed a significant increase of AAV.hCRB1 in AAV.hCRB1-treated and AAV.hCRB2 in AAV.hCRB2-treated organoids in the RPE of the *CRB1* patient-derived retinal organoids (Figures S5B and S5C).

Differential gene expression followed by GO term analysis of AAV.hCRB1 treatment compared with untreated P128 retinal organoids in MGCs revealed differences related to the endosomal system, cell-cell adhesion, and protein or receptor binding (Figure 5A). Also for AAV.hCRB2 treatment similar terms were observed to be statistically significant in P128 (Figure 5C). Next, these GO terms and differentially expressed genes were compared with the ones observed when contrasting

(C) Immunohistochemical image of CRB2 in untreated and AAV.hCRB2-treated *CRB1* patient-derived retinal organoid at the OLM. Arrows indicate overexpression of CRB2 in photoreceptor cells in AAV.hCRB2-treated retinal organoids.

(D–F) Representative immunohistochemical images of (D) untreated, (E) AAV.hCRB1-, and (F) AAV.hCRB2-treated *CRB1* patient retinal organoids stained with MUPP1 (red) at DD210.

(G) Quantification of the number of photoreceptor nuclei in a row (from left to right:  $p = 0.000$ ,  $p = 0.000$ ,  $p = 0.039$ ,  $p = 0.046$ ,  $p = 0.046$ ), (H) ONL thickness ( $p = 0.001$ ,  $p = 0.001$ ,  $p = 0.923$ ,  $p = 0.757$ ,  $p = 0.243$ ), and (I) the number of photoreceptor nuclei above the OLM ( $p = 0.116$ ,  $p = 0.034$ ,  $p = 0.717$ ,  $p = 0.730$ ) in three *CRB1* patients and three isogenic control organoids with and without concomitant treatment of AAV.GFP pooled. Scale bars, 50  $\mu\text{m}$ . Each datapoint in the graph represent individual organoids, of which an average has been taken of at least three representative images. The standard error of mean (SEM) is derived from these averages. Number of individual organoids per condition: *Crb1* patients (P116, P117, and P128 pooled) treated with AAV.hCRB1 ( $n = 34$ ), AAV.hCRB2 ( $n = 33$ ), untreated and GFP-treated ( $n = 32$ ), and isogenic controls (ISO-02 P116, ISO-03 P116, ISO-P128 pooled) treated with AAV.hCRB1 ( $n = 14$ ), AAV.hCRB2 ( $n = 10$ ), and untreated and GFP-treated ( $n = 24$ ) independent organoid from two different differentiation rounds. Statistical tests: generalized linear mixed models with  $*p < 0.05$ ,  $**p < 0.01$ ,  $***p < 0.001$ . Related to Figure S4.



**Figure 5. scRNA-seq of *CRB1* patient-derived retinal organoid treated with AAV.hCRB1 or AAV.hCRB2 restores transcriptional effect on the endosomal system**

(A and C) Gene ontology (GO) analysis of differentially expressed markers contrasting untreated with (A) AAV.hCRB1- or (C) AAV.hCRB2-treated P128 in MGCs clustered in groups with similar terms in the same color.

(B and D) All significantly differentially expressed markers in terms related to the endosomal system after treatment with (B) AAV.hCRB1 or (D) AAV.hCRB2. All markers present in (B and D) are also statistically significant for P128 compared with ISO-P128. Number of independent organoids used: P128, n = 6; P128+ AAV.hCRB1, n = 5; P128+ AAV.hCRB2, n = 5, from one differentiation round equally divided into three separate sequencing rounds. Related to [Figure S5](#) and [Table S5](#).

P128 with ISO-P128. Overlapping differentially expressed genes associated with the endosomal system show that, after AAV.hCRB1 treatment, the expression levels from the patient retinal organoids are similar to levels of the isogenic control ([Figure 5B](#)). Moreover, after AAV.hCRB2 treatment the genes associated with the endosomal system appeared to be restored as well in MGCs ([Figure 5D](#)).

Similar comparisons were performed for the other two *CRB1* patient-derived retinal organoids. For P116 and P117, statistically significant GO terms related to the endosomal system were also observed after AAV.hCRB1 and AAV.hCRB2 treatment ([Figures S5D, S5E, S5G, S5I, and S5K](#)). When analyzing in more detail the differentially expressed genes associated with the endosomal system in AAV.hCRB1-treated P116 organoids, transcript levels

seemed to be restored to isogenic control levels in MGCs ([Figure S5J](#)) as well as in rod photoreceptor cells ([Figure S5L](#)). The genes for ISO-P116, indicated with a dashed line box around them, were not statistically significantly different from P116, but the genes after AAV.hCRB treatment are similarly expressed as the average expression in ISO-P116 ([Figure S5J](#)). Similar results were observed for genes associated with the endosomal system in AAV.hCRB2-treated organoids ([Figure S5I](#)). Moreover, after both AAV.hCRB1 and AAV.hCRB2 treatment in MGCs of P117, we observed that the expression levels changed in a similar direction ([Figure S5F](#)).

In summary, AAV.hCRB1 as well as AAV.hCRB2 treatment on *CRB1* patient-derived retinal organoids restores gene expression related to the endosomal system back to isogenic control levels.



## DISCUSSION

In this manuscript we (1) showed diminished levels of variant CRB1 protein in *CRB1* RP patient retinal organoids that harbor missense mutations, (2) demonstrate moderate loss of photoreceptors in *CRB1* patient-derived retinal organoids at DD210, (3) detect transcriptional differences suggesting changes within the endosomal system in *CRB1* patient compared with isogenic control organoids, (4) show that AAV5.CMV.GFP efficiently transduced MGCs in addition to photoreceptors and RPE at DD120, and (5) observe a partially restored phenotype after AAV.h*CRB1*- or AAV.h*CRB2*-mediated gene therapy in *CRB1* patient-derived retinal organoids.

*CRB1* patients' clinical and genetic characteristics were described previously in detail in a prospective natural history study on 22 patients (Nguyen et al., 2022). The P116 retinal organoids were derived from skin fibroblasts from a patient with a first diagnosis RP and discontinuous OLM and ellipsoid zone (EZ) in parafovea and perifovea on spectral domain optical coherence tomography (see Tables S1 and S2 in Nguyen et al. [2022]). The P117 retinal organoids were derived from a patient who experienced mild RP with at first diagnosis loss of visual acuity and continuous OLM and EZ in parafovea and perifovea. The P128 retinal organoids were derived from a patient with at first diagnosis RP, nyctalopia, and discontinuous OLM and EZ in parafovea and perifovea. Here, the *CRB1* patient-derived retinal organoids were compared with corresponding isogenic controls at DD210. We show a decreased number of photoreceptor nuclei in a row and a reduced ONL thickness in the *CRB1* patient-derived retinal organoids compared with the isogenic controls. Decreased levels of variant CRB1 at the OLM of *CRB1* patient-derived retinal organoids might be associated with increased protrusion of photoreceptor cell bodies into the cell culture medium at DD180 (Quinn et al., 2019b) and thinning of the photoreceptor ONL at DD210 (current article). This process is similar to the complete loss of CRB1 at the OLM in *Crb1* mouse retina which results in protrusion of photoreceptor cell bodies into the subretinal space (van de Pavert et al., 2004, van de Pavert et al., 2007a, 2007b) or the loss of CRB2 at the OLM in *Crb2* mouse retina (Alves et al., 2013).

Moreover, it was shown that CRB1 and CRB2 are present at the OLM of both photoreceptor and MGCs in iPSC-derived retinal organoids (Quinn et al., 2019b). Our scRNA-seq data confirm on the transcriptome, with more *CRB1* expression in MGCs than in photoreceptor cells and more *CRB2* expression in photoreceptor cells than in MGCs. The rather high expression levels of *CRB1* and low levels of *CRB2* in MGCs might be related to the phenotype variation, since mutations in *CRB1* may cause either early

onset RP or LCA. Variable low levels of *CRB2* transcripts in MGCs of *CRB1* patients may be involved in the severity of the phenotype. Such a hypothesis would be in strong correlation with our previous studies in mice, which suggests a modifying role for *CRB2* in *CRB1*-related dystrophies (Buck et al., 2021; Pellissier et al., 2014b, 2015; Quinn et al., 2018, 2019a).

*CRB1* variant protein at the OLM was strongly diminished in *CRB1* patient-derived retinal organoids, while *CRB1* expression levels remained similar. In contrast, CRB core complex members as well as the FERM proteins remained at the OLM and similar expression levels were observed. This indicates that a variant CRB1 protein is produced but it does not localize to or maintain its expected location at the OLM. The endolysosomal system is required for transport of CRB1 to the OLM but also for recycling of endocytosed CRB1 from the early endosome to the OLM and the transport into degradative vesicles. *Drosophila* studies show that CRB trafficking is mediated by transport along microtubules by Rab11- and retromer-containing endosomes (Aguilar-Aragon et al., 2020; Pocha et al., 2011). In addition, in *Drosophila* salivary gland cells Crumbs maintains the active pool of Rab proteins at the apical domain, which is essential for maintaining the organization of the apical membrane and efficient apical secretion (Lattner et al., 2019). The scRNA-seq GO data shown here suggest an aberrant endosomal pathway specifically in MGCs and rods of *CRB1* patient-derived retinal organoids. Dysregulation of CRB1 at the OLM can thus cause changes in the endosomal system. Endosomal recycling is pivotal for maintenance of neuronal health, and defects in its function results in human neurodegenerative disorders (Saitoh, 2022; Tang et al., 2015). We hypothesize that the reduced levels of variant CRB1 at the OLM are caused by disturbed variant CRB1 protein transport to the OLM, or disturbed endosomal recycling of variant CRB1 between OLM and the early endosome, or increased variant CRB1 degradation in the retinal organoids. Preliminary studies suggest changes in the recycling endosome and in degradative vesicles (T.M.B. and J.W., unpublished data). In analogy to the roles of the Crumbs protein in *Drosophila* salivary glands (Lattner et al., 2019), in future studies we will examine the putative role for CRB1 in the maintenance of an active pool of RAB11 and VPS35 (retromer) recycling endosome proteins at the OLM.

Previously, we described an improved phenotype after AAV.h*CRB2* treatment in *Crb* mutant mouse models (Buck et al., 2021; Pellissier et al., 2015). Here, we investigated whether we could observe an improved *CRB1*-RP phenotype after AAV-mediated gene augmentation therapy in *CRB1* patient-derived retinal organoids. Proof-of-concept for developing gene therapy in retinal organoids for CRX-LCA has been described, where AAVs were used to alleviate



the phenotype observed in *CRX* mutant retinal organoids (Kruczek et al., 2021). In addition, AAV-mediated gene augmentation of RP2 knockout retinal organoids prevents ONL thinning and restored rhodopsin expression (Lane et al., 2020). In this article, using AAV.h*CRB1* and AAV.h*CRB2* gene augmentation therapy, a partially restored phenotype was observed in *CRB1* patient-derived retinal organoids. The number of photoreceptor nuclei in a row and ONL thickness were significantly improved after AAV.h*CRB* treatment when analyzed at DD210, showing the long-term effects of the gene augmentation therapy. Moreover, neither positive nor negative effects were observed when treating isogenic controls with AAV.h*CRB1* or AAV.h*CRB2*. Furthermore, the infection of AAV.h*CRB1* on the *CRB1*<sup>KO</sup> retinal organoids shows localization of recombinant CRB1 protein at the OLM. Whereas the recombinant CRB1 protein localizes merely at the OLM, we also detected CRB1 protein around the OLM as detected previously in first trimester human fetal retina and DD120 immature wild-type retinal organoids (Quinn et al., 2019b). The imprecise localization of recombinant CRB1 is potentially related to a partial restoration of CRB1-positive recycling endosomal vesicles at the OLM and is the subject of future studies. To our knowledge, this is the first time that an improved phenotype after AAV.h*CRB* gene augmentation in *CRB1* patient-derived retinal organoids has been observed.

In conclusion, we demonstrate in *CRB1* patient-derived retinal organoids a moderate loss of photoreceptor nuclei in a row, strongly reduced levels of CRB1 variant protein with unaffected *CRB1* transcript levels, and a dysregulated molecular gene profiling phenotype of MGCs and rod photoreceptor cells, suggesting an aberrant endosomal system. Moreover, using AAV-mediated gene augmentation therapy approaches we have improved the histological and transcriptional retinal phenotype in *CRB1* patient-derived retinal organoids. These data provide essential information for future gene therapy approaches for patients with mutations in the *CRB1* gene.

## EXPERIMENTAL PROCEDURES

### Resource availability

#### Corresponding author

Further information and requests for resources and reagents should be directed to and will be fulfilled by the corresponding author, Jan Wijnholds ([j.wijnholds@lumc.nl](mailto:j.wijnholds@lumc.nl)).

#### Materials availability

Materials and additional details can be made available from the corresponding author upon reasonable request.

#### Data and code availability

scRNA-seq data are available at the NCBI Gene Expression Omnibus database (GEO: GSE212582).

### Cell culture and retinal organoid differentiation

The following hiPSC lines were used for organoid differentiation: three *CRB1* RP patient-derived lines and one control (LUMC0116iCRB09, LUMC0117iCRB01, LUMC0128iCRB01, LUMC0004iCTRL10 [Quinn et al., 2019b]), and three isogenic controls of the *CRB1* patient-derived lines (LUMC0116iCRB-ISO02, LUMC0116iCRB-ISO03, LUMC0128iCRB-ISO01) (Figure S1; Table S1). hiPSC lines were derived from skin fibroblast using polycistronic lentiviral vectors (Warlich et al., 2011).

hiPSCs were maintained on Matrigel-coated plates in mTeSR plus medium and passaged mechanically using gentle cell dissociation reagent (STEMCELL Technologies). Retinal organoid differentiation was carried out as reported previously with some modifications (experimental procedures in supplemental information) (Quinn et al., 2019b; Zhong et al., 2014). Retinal organoids were collected at DD180 or DD210 for immunohistochemical analysis; a list of all primary antibodies used for immunofluorescent staining is provided in (Table S2). At least three different differentiation batches were analyzed to verify disease phenotypes.

### AAV transduction of hiPSC-derived retinal organoids

Two to three retinal organoids were plated in a 96-well agarose-coated plate and were infected with AAV in 50  $\mu$ L RLM2 and incubated for 8 h at 5% CO<sub>2</sub> at 37°C. After this, the wells were topped up to 200  $\mu$ L with RLM2. The next morning, treated organoids were transferred to a 24-well plate and cultured for at least 3 weeks or until the desired differentiation day. AAV5.CMV.GFP and AAV2.CMV.GFP (105530; Addgene) were used at titers of  $1 \times 10^{10}$ ,  $3.3 \times 10^{10}$ ,  $6.6 \times 10^{10}$ , or  $10 \times 10^{10}$  gc. AAV5.CMV.min.h*CRB1* and AAV5.CMV.h*CRB2* (HORAMA) were used at a titer of  $3.3 \times 10^{10}$  gc.

### Quantification and statistical analysis

Magnification images (40 $\times$ ) were manually quantified using Fiji ImageJ (ImageJ 1.53f51). At least four organoids per condition with three to six representative images of each organoid were used for quantification. Three regions in each image were manually analyzed for the number of photoreceptor nuclei in a row in the ONL, the number of photoreceptor nuclei above the OLM, retinal thickness, INL thickness, and ONL thickness. Quantifications were performed independently by at least two researchers without the knowledge of genotype or treatment. For the MUPP1 quantifications, a ROI was drawn at the OLM and the average intensity was measured using ImageJ. All datapoints measured were averaged per organoid and plotted in the graph; so that each point is one organoid. No normalization of the values was performed. Data were either presented per 100  $\mu$ m retinal length or per field of view. Data presentation and statistical analysis were performed using GraphPad Prism version 8 (GraphPad Software) and IBM SPSS statistics (version 25), respectively. For statistical analysis, all individual values per image were used. A generalized linear mixed model with treatment (and patient) as a fixed effect was performed on all quantification parameters; the statistical test took into account that multiple *CRB1* patients were merged by introducing a random intercept per patient. Data are presented as mean per organoid  $\pm$  standard error of the mean. Significance is indicated in graphs as \* $p < 0.05$ , \*\* $p < 0.01$ , and \*\*\* $p < 0.001$ .



## scRNA-seq

Retinal organoids were dissociated using an adapted protocol from the Papain Dissociation Kit (Worthington, I-LK 03150). Analysis and processing of single-cell transcriptomics using Seurat is detailed in [experimental procedures](#) in [supplemental information](#).

## SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.stemcr.2023.03.014>.

## AUTHOR CONTRIBUTIONS

Conceptualization, N.B. and J.W.; software, N.B., I.M., S.B., and H.M.; formal analysis, N.B.; investigation, N.B., X.L., C.A.A., and T.M.B.; resources, C.F. and C.H.A.; data curation, N.B. and I.M.; writing – original draft, N.B.; writing – review & editing, N.B. and J.W.; visualization, N.B.; project administration, J.W.; funding acquisition, J.W.

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## CONFLICT OF INTERESTS

The LUMC is the holder of patent application PCT/NL2014/050549, which describes the potential clinical use of CRB2; J.W. is listed as inventor on this patent, and J.W. is an employee of the LUMC. Horama (now known as Coave Therapeutics) partially sponsored this study. N.B., X.L., C.A.A., and J.W. received funding from Horama.

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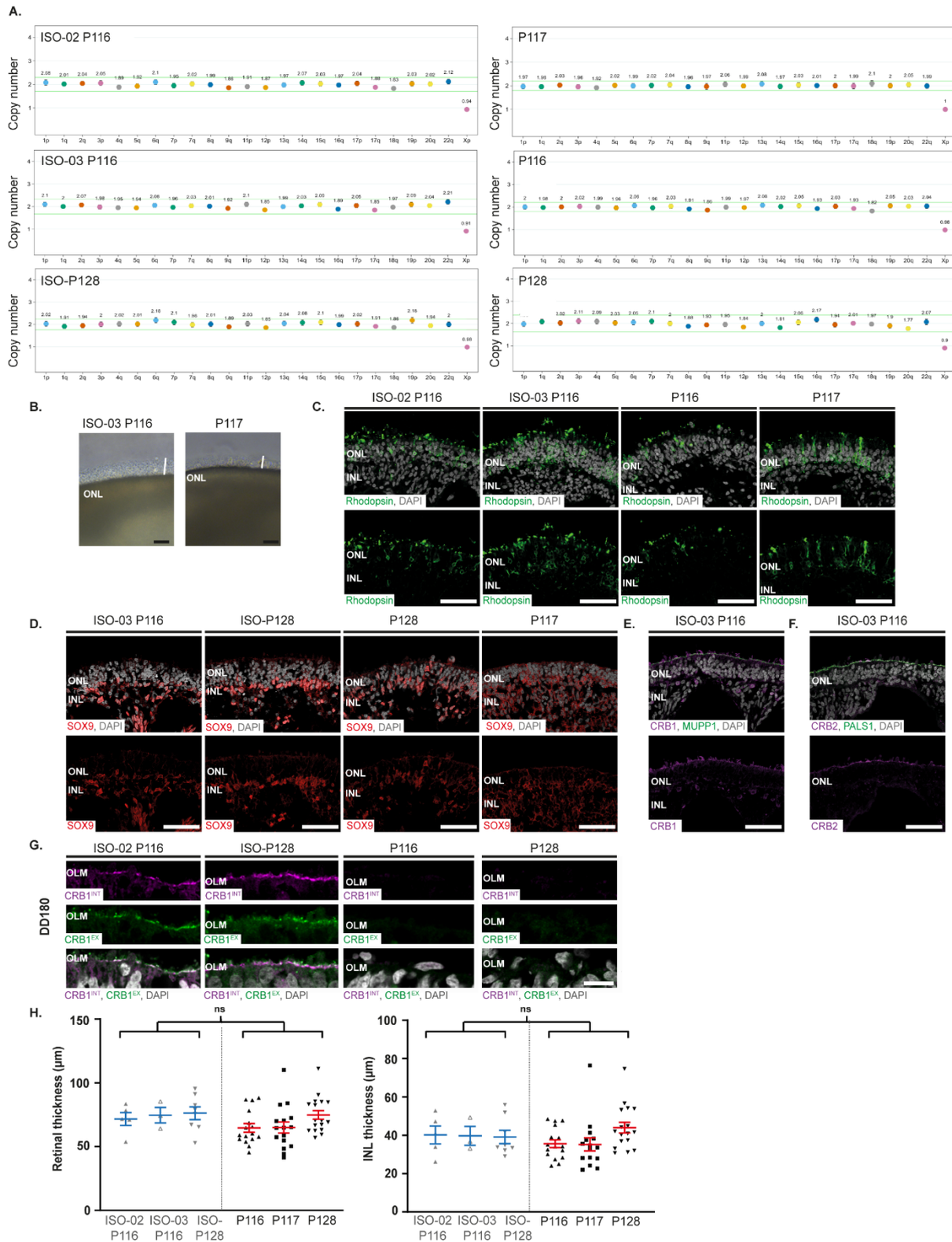
**Stem Cell Reports, Volume 18**

## **Supplemental Information**

### **AAV-mediated gene augmentation therapy of *CRB1* patient-derived retinal organoids restores the histological and transcriptional retinal phenotype**

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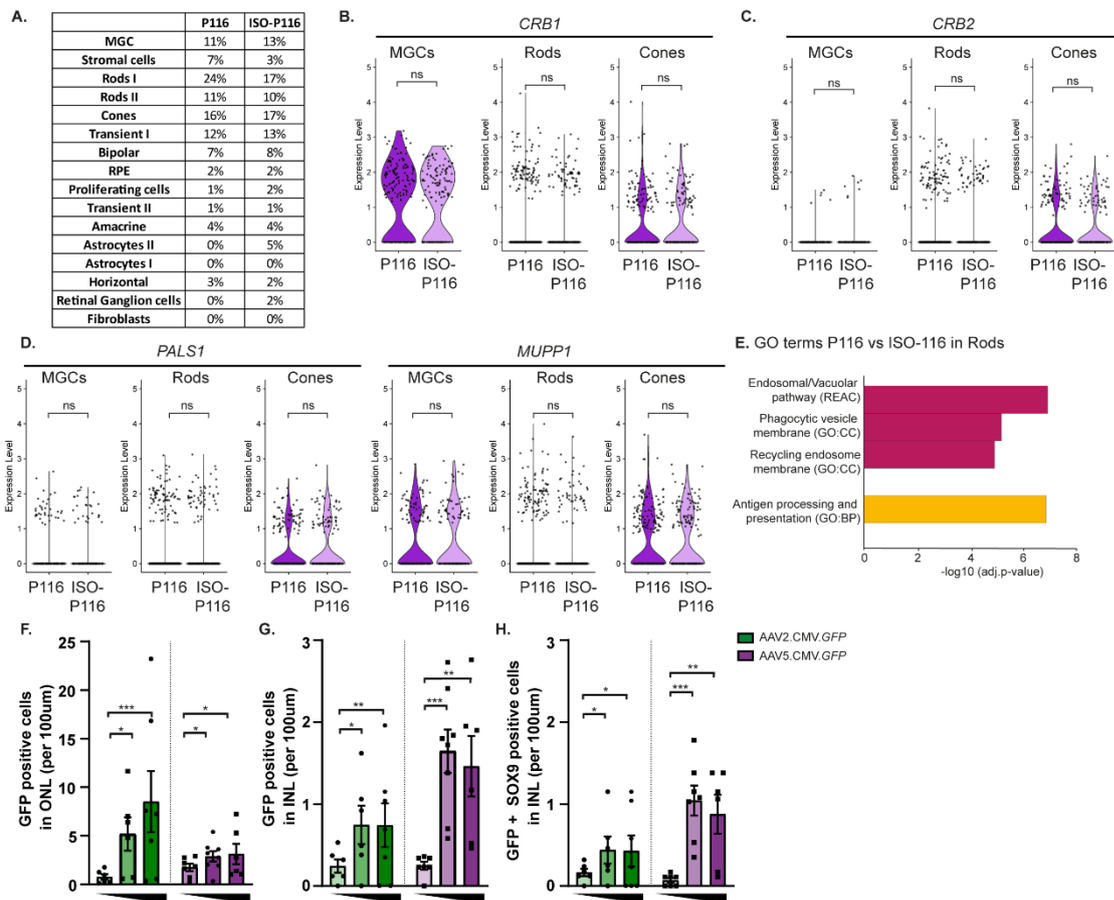
## Supplemental Figures and Legends



03 P116. (E, F) Representative immunohistochemical images of (E) CRB1 (magenta) co-localized with MUPP1 (green) and of (F) CRB2 (magenta) co-localized with PALS1 (green) in ISO-03 P116. (G) Representative immunohistochemical images of CRB1<sup>EX</sup> (green) and CRB1<sup>INT</sup> (magenta) in *CRB1* patient-derived retinal organoids compared to isogenic controls at DD180. (H) Quantitative analysis of the total retinal thickness ( $p=0.158$ ) and INL thickness ( $p=0.696$ ) per field of view in *CRB1* patient derived and isogenic control retinal organoids. Each datapoint in the graph represent individual organoids, of which an average has been taken of at least 3 representative images. The standard error of mean (SEM) is derived from these averages. Number of individual organoids per condition and differentiation round: P116  $n=16$ , P117  $n=15$ , P128  $n=17$  from four independent organoid batches, ISO-P128  $n=8$  from three independent organoid batches, ISO-02 P116  $n=5$  and ISO-03 P116  $n=5$  from two independent organoid batches. Scalebar = (C-F) 50 $\mu$ m, (G) 10 $\mu$ m.



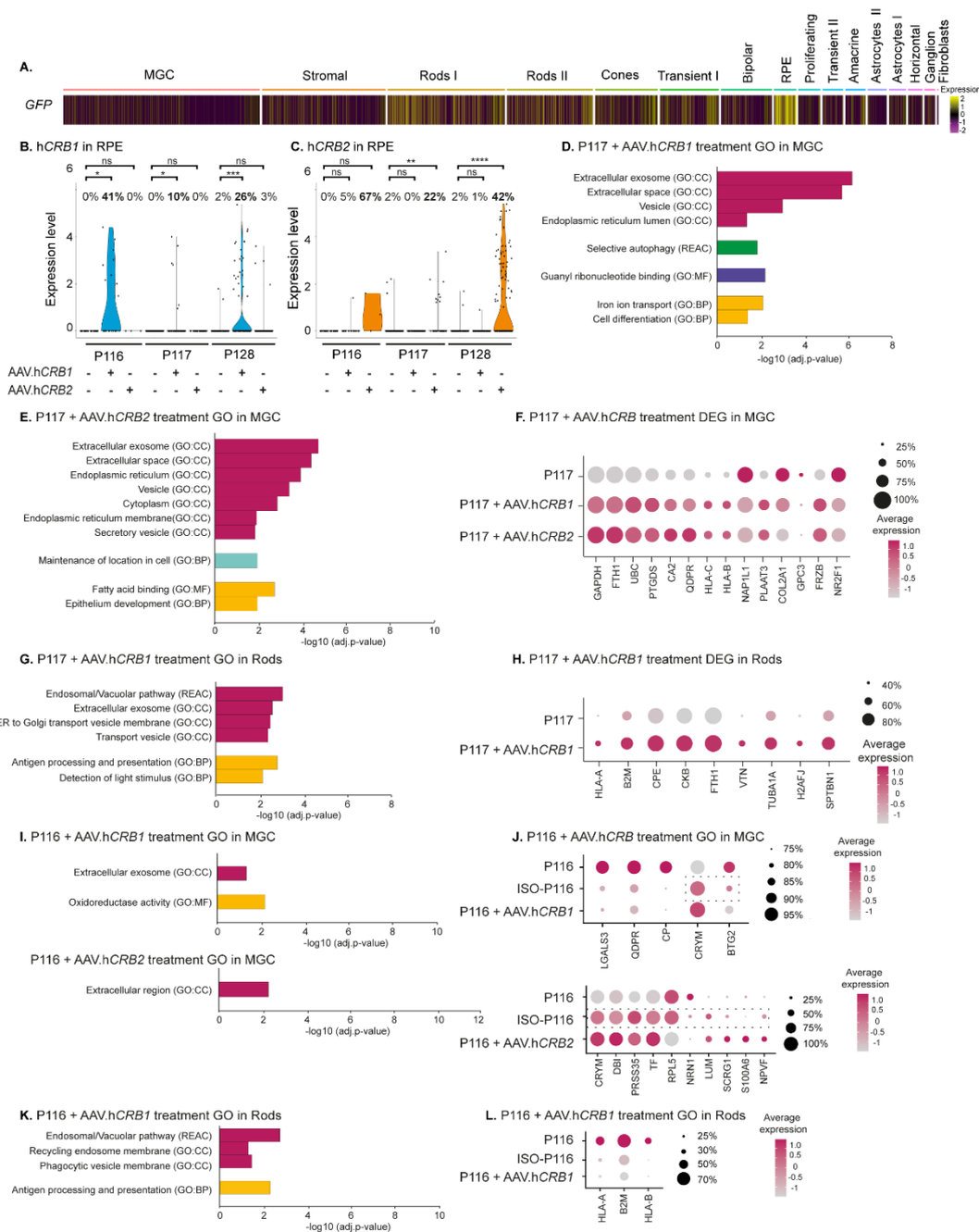
p=0.41, p=0.17, p=0.59), rods (p=0.018, p=0.89, p=0.53) and cones (p=0.77, p=0.79, p=0.24) in P128 vs ISO-P128. Number of independent organoids used: ISO-P128  $n=6$ , and P128  $n=6$  from one differentiation round equally divided into three separate sequencing rounds.



**Figure S3: scRNA-seq analysis comparing ISO-P116 and P116 retinal organoids.** Related to Figure 2. (A) Table showing all retinal cell types equally present in the retinal organoids. (B) Violin plots of *CRB1* and (C) *CRB2* expression levels specifically in MGCs (p=0.78, p=0.28), rods (p=0.37, p=0.88), and cones (p=0.51, p=0.96). (D) Violin plots of canonical core CRB complex members *PALS1*, and *MUPP1* levels in (p=0.49, p=0.98), rods (p=0.36, p=0.61), and cones (p=0.41, p=0.86). (E) Gene ontology (GO) analysis of differentially expressed markers specifically in rods clustered with similar terms in the same colour. Number of independent organoids used: ISO-P116  $n=4$ , and P116  $n=4$  from one differentiation and sequencing round. (F, G, H) Quantification of AAV2.CMV.GFP and AAV5.CMV.GFP retinal organoids at DD120 with three different titre concentrations:  $1 \times 10^{10}$ ,  $6.6 \times 10^{10}$ , and  $10 \times 10^{10}$  gc (genome copies) in the (F) ONL, (G) INL, and (H) GFP positive MGC in the INL per 100µm. Each datapoint in the graph represent individual organoids, of which an average has been taken of at least 3 representative images. The standard error of mean (SEM) is derived from these averages. Number of organoids per condition: for AAV2.CMV.GFP  $1 \times 10^{10}$   $n=5$ ,  $6.6 \times 10^{10}$   $n=6$ , and  $10 \times 10^{10}$   $n=7$ , and for AAV5.CMV.GFP  $1 \times 10^{10}$   $n=7$ ,  $6.6 \times 10^{10}$   $n=8$ , and  $10 \times 10^{10}$   $n=6$  individual organoids from two independent differentiation rounds. Statistical analysis: generalized linear mixed models with  $p < 0.05$  (\*),  $p < 0.01$  (\*\*), and  $p < 0.001$  (\*\*\*).



immunohistochemical images of anti-CRB1<sup>EX</sup> and anti-CRB1<sup>INT</sup> showing localization in the RPE after AAV.hCRB1 treatment in *CRB1* patient-derived retinal organoid. (G) Immunohistochemical image of anti-CRB1<sup>INT</sup> showing localization in the OLM after AAV.hCRB1 treatment in *CRB1*<sup>KO</sup> retinal organoid. (H, I) Representative immunohistochemical images of CRB2 at the OLM (H) and RPE (I) after AAV.hCRB2 treatment in *CRB1* patient-derived retinal organoids. (J) Quantification of the average MUPP1 fluorescence intensity at the OLM of DD210 *CRB1* patient derived retinal organoids (P116, P117, P128 pooled) treated with AAV.hCRB. (K) Representative immunohistochemical images of PALS1 at the OLM with and without AAV.hCRB treatment in a *CRB1* patient-derived retinal organoid. (L) Quantification of the retinal thickness ( $p=0.008$ ,  $p=0.082$ ,  $p=0.993$ ,  $p=0.981$ ,  $p=0.981$ ) and (M) the INL thickness ( $p=0.139$ ,  $p=0.632$ ,  $p=0.958$ ,  $p=0.195$ ,  $p=0.707$ ) per field of view of *CRB1* patient and isogenic control retinal organoids at DD210. (N) SDS-PAGE gel of AAV.hCRB1 and AAV.hCRB2 showing no contamination in the AAV preparation. Scalebar = 50 $\mu$ m. Each datapoint in the graph represent individual organoids, of which an average has been taken of at least 3 representative images. The standard error of mean (SEM) is derived from these averages. Number of individual organoids per condition at DD180 P117 treated with AAV.hCRB1  $n=13$ , AAV.hCRB2  $n=12$ , untreated  $n=14$  from two different differentiation rounds, control organoids treated with AAV.hCRB1  $n=4$ , AAV.hCRB2  $n=3$ , and untreated  $n=2$  from one differentiation round. And at DD210 *CRB1* patient-derived retinal organoids (P116, P117, P128 pooled) treated with AAV.hCRB1  $n=34$ , AAV.hCRB2  $n=33$ , untreated  $n=32$ , and isogenic controls (ISO-02 P116, ISO-03 P116, ISO-P128 pooled) treated with AAV.hCRB1  $n=14$ , AAV.hCRB2  $n=10$ , and untreated  $n=24$  independent organoid from two different differentiation rounds. Statistical tests: generalized linear mixed models with  $p<0.05$  (\*),  $p<0.01$  (\*\*), and  $p<0.001$  (\*\*\*)



**Figure S5: Single cell RNA-sequencing of *CRB1* patient-derived retinal organoid treated with AAV.*hCRB1* or AAV.*hCRB2* restores transcriptional effect on the endosomal system.** Related to Figure 5. (A) Heatmap showing *AAV.GFP* expression in the clusters, mainly transducing MGC, photoreceptor cells, and RPE. (B, C) Violin plot of transcript expression of (B) AAV.*hCRB1* (from left to right:  $p=0.018$ ,  $p=0.41$ ,  $p=0.041$ ,  $p=0.078$ ,  $p<0.001$ ,  $p=0.54$ ) or (C) AAV.*hCRB2* ( $p=0.5$ ,  $p=0.1$ ,  $p=0.38$ ,  $p=0.0037$ ,  $p=0.53$ ,  $p<0.00001$ ) in the RPE of AAV.*hCRB* treated retinal organoids. (D, E) Gene ontology (GO) analysis of differentially expressed markers contrasting untreated with (D) AAV.*hCRB1* or (E) AAV.*hCRB2* treated P117 in MGCs clustered in groups with similar terms in the same colour. (F) All significantly differentially expressed markers in terms related to the endosomal system after treatment with AAV.*hCRB1* or AAV.*hCRB2* in MGC. (G, H) P117 treated with AAV.*hCRB1* in rods showing gene ontology (G) and all terms related to the endosomal system (H). (I)

GO of P116 treatment with AAV.hCRB in MGC and (J) dot plot of terms related to the endosomal system after treatment. The data in the box with a dashed line are not statistically significant different from P116, all the other data points (in F, H, J, and L) are statistically significant different from untreated patient derived retinal organoids. Number of independent organoids used: P116  $n=4$ , P116 + AAV.hCRB1  $n=3$ , P116 + AAV.hCRB2  $n=3$  from one differentiation and sequencing round, and P117  $n=5$ , P117 + AAV.hCRB1  $n=5$ , P117 + AAV.hCRB2  $n=5$  from one differentiation round equally divided into three separate sequencing rounds.

### Supplemental Tables

**Table S1. hiPSC line information.** Related to all figures.

Previously published line name	Description	Gender
LUMC04iCTRL10 (Quinn et al., 2019)	Control iPSC line	Male
CRB1 <sup>KO</sup> LUMC04iCTRL10	CRB1 <sup>KO</sup> line was derived from LUMC04iCTRL10; it has a stop codon in the second exon of CRB1 (Boon N et al Wijnholds J, unpublished data). Only used for proof of recombinant CRB1 protein expression after AAV.hCRB1 treatment.	Male
LUMC0116iCRB09 (Quinn et al., 2019)	<u>P116</u> = Allele 1 and 2: homozygous c.3122T>C--> p.(Met1041Thr)	Male
iso02LUMC0116iCRB09	<u>ISO-02 P116</u> = Allele 1: c.3122T>C gene corrected to c.3120C>G. Allele 2: c.3122T>C. p.(Met1041Thr)	Male
iso03LUMC0116iCRB09	<u>ISO-03 P116</u> = Homozygous c.3122T>C gene corrected to c.3120C>G	Male
LUMC0117iCRB01 (Quinn et al., 2019)	<u>P117</u> = Allele 1: c.1892A>G (p.Tyr631Cys). Allele 2: c.2911G>T (p.(Glu995*))	Male
LUMC0128iCRB01 (Quinn et al., 2019)	<u>P128</u> = Allele 1: c.2843G>A --> p.(Cys948Tyr). Allele 2: c.3122T>C --> p.(Met1041Thr)	Male
iso02LUMC0128iCRB01	<u>ISO-P128</u> = Allele 1: c.2843G>A. p.(Cys948Tyr). Allele 2: c.3122T>C gene corrected to c.3120C>G	Male

**Table S2. List of primary antibodies used in the study.** Related to all figures.

Antigen	Dilution	Source	Identifier
CRB1 (intracellular domain – used if not otherwise specified)	1:200	Homemade	NA
CRB1 (extracellular domain)	1:200	Abnova	H00023418-A01
CRB2	1:200	Homemade	NA
CRALBP	1:200	Abcam	ab15051
MUPP1	1:200	BD Biosciences	M98820

PALS1	1:200	Homemade	NA
OTX2	1:200	Proteintech	13497-1-AP
Rhodopsin	1:200	Sigma	SAB4502636
Glutamine synthetase (GS)	1:250	BD Biosciences	610518
SOX9	1:250	Millipore	AB5535

**Table S3. Top differentially expressed genes per cluster.** Related to Figure 2. Statistically significant log2 fold changes of the expression level of differentially expressed genes in the cluster (pct1) comparing with the remaining clusters (pct2). With the defined cell type per cluster.

**Table S4. Differentially expressed genes markers and gene ontology terms contrasting P128 and ISO-P128, and P116 and ISO-P116.** Related to Figure 2, S3A-E. Log2 fold changes of the expression level of statistically significant expressed genes in *CRB1* patient-derived retinal organoids comparing with the isogenic control in MGCs or in rod photoreceptor cells and the associated gene ontology terms.

**Table S5. DEG markers and gene ontology terms comparing untreated and AAV.hCRB treated CRB1 patient derived retinal organoids.** Related to Figure 5, S5. Log2 fold changes of the expression level of statistically significant expressed genes in AAV.hCRB treated *CRB1* patient-derived retinal organoids comparing with the untreated and the associated gene ontology terms.

### Supplemental experimental procedures

#### Cell culture and retinal organoid differentiation

Human induced pluripotent stem cells (hiPSC) were maintained on Matrigel coated plates in mTeSR plus medium (STEMCELL Technologies) and passaged mechanically. Retinal organoid differentiation was carried out as previously reported with some modifications (Quinn et al., 2019; Zhong et al., 2014). Confluent hiPSCs were collected and incubated with ( $\pm$ )blebbistatin in mTeSR medium in micro-mold spheroids (Z764000-6EA, Merck) over night. Then, medium was transitioned to Neural Induction Medium 1 (NIM1) using mTeSR/NIM1 (3:1), then (1:1), and finally (0:1) over three days to form embryoid bodies (EBs). After 1 week, EBs were plated onto Matrigel-coated wells with daily NIM-1 medium change till DD15 and daily change of NIM-2 starting at DD16. Between DD20 and latest DD28, neuroepithelial structures were selected and flushed from the Matrigel plates using a P1000 pipet and kept in floating culture in agarose coated plates from this point onwards. After selecting the best-looking structures, all structures were flushed from the Matrigel plates and kept in floating culture to increase the yield of obtained organoids. Then, typically from DD40 until DD100, good retinal organoid structures were selected and placed individually in a 48 well plate. Brain and other non-retinal structures were removed as well between this time period. Daily medium change of NIM-2 is used till DD34, then typically three times a week Retinal Lamination Medium 1 (RLM-1) was used

from DD35 to DD63. Then, RLM-1 + 1 $\mu$ M retinoic acid until DD84, followed by RLM-2 + 0.5 $\mu$ M retinoic acid, and RLM-2 from DD120 was used for the rest of the culture.

### **Immunohistochemical analysis**

Organoids were collected at DD180 or DD210 for immunohistochemical analysis. Organoids were fixed with 4% paraformaldehyde in PBS for 20 minutes at RT, briefly washed with PBS and subsequently cryo-protected with 15% and 30% sucrose in PBS until organoids sunk to the bottom of the well. Organoids were embedded in Tissue-Tek O.C.T. Compound (Sakura, Finetek), thereafter 8  $\mu$ m cryosections were made with a Leica CM1900 cryostat (Leica Microsystems) and stored in the freezer.

For immunohistochemistry, the sections were blocked for 1h at RT in 10% normal goat serum, 0.4% Triton X-100, and 1% bovine serum albumin in PBS. Primary antibodies were incubated overnight at 4°C or for at least 3h at RT with 0.3% normal goat serum, 0.4% Triton X-100, 1% BSA and appropriate primary antibody concentration (Table S2). Then, slides were washed for two times 15 minutes in PBS and subsequently incubated for 1h at RT with fluorescent-labelled secondary antibody in 0.1% goat serum in PBS. Nuclei were counterstained with DAPI and mounted in Vectashield Hardset mounting medium (H1800, Vector laboratories, Burlingame, USA). Sections were imaged on a Leica TCS SP8 confocal microscope and images were processed with Leica Application suite X (v3.7.0.20979).

### **RNA isolation, cDNA synthesis, and qPCR analysis**

RNA was isolated from DD210 retinal organoids of P116, P117, P128, ISO-P128, and ISO-02 P116 using TRIZOL reagent (Gibco Life Technologies) according to the manufacturer manual. The isolated RNA was dissolved in 20 $\mu$ l RNase-free water. 0.5 $\mu$ g of total RNA was reverse transcribed into first-strand cDNA using QuantiTect Reverse Transcription Kit (205311, QIAGEN) in a total reaction volume of 20 $\mu$ l. From all cDNA samples, a 1 in 20 dilution was made and used for qPCR analysis.

Two different exon-spanning primer pairs were designed at the 5' end of the *CRB1-B* gene giving rise to an amplicon of 70 to 120bp (FW1: TGTTTGGAGCCAGGACACAT, REV1: ACGTCTTCTTCGCAGTGGAT and FW2: GAGCCAGGACACATGGTTTTTC, REV2: TTCCCAGGCAAGTTCTCACA). Real-time qPCR was based on the monitoring of SYBR Green I dye fluorescence on a CFX Connect Real-Time System (BioRad). The qPCR conditions were as follows: 5 $\mu$ l SYBR green PCR 2x master mix (4913914001, Merck), 0.2 $\mu$ l of 10 $\mu$ M FW and REV primers, and 5 $\mu$ l of the diluted cDNA. qPCR machine started with a melting step at 95°C for 10min, followed by 40 cycles of 95°C for 15 seconds and an annealing at 60°C for one minute. At the end of the PCR run, a dissociation curve was determined by ramping the temperature of the sample from 60 to 95°C while continuously collecting fluorescence data. MQ water controls were included for each primer pair to check for any significant levels of contaminants. The following two reference genes were used: glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and elongation factor 1a (EF1A), previously described by (Pellissier et al., 2014).

qPCR was performed on both primer pairs with at least three individual DD210 retinal organoids of P116, P117, P128, ISO-P128, and ISO-02 P116 cDNA and human adult retina cDNA (Marathon-ready; Clontech). *CRB1-B* was detected in adult human retina cDNA but was below detection level in the patient-derived and isogenic control retinal organoids at DD210.

### **Single cell RNA sequencing**

#### *Retinal organoid dissociation*

Retinal organoids were dissociated using an adapted protocol from the Papain Dissociation kit (Worthington, I-LK 03150). In short, single retinal organoids were selected and cut into small pieces to remove excess RPE or non-neural tissue as much as possible and placed on a 48 well plate with 500µl dissociation solution (20 units/ml papain and 0.005% DNase). These were incubated for 30 minutes on a shaker in the incubator, then the organoids were triturated using a 1mL pipettor to dissociate the tissue. The plate was placed back for 15-20 minutes on the shaker in the incubator, then again triturated, this was repeated until a single cell suspension was obtained. 500µl albumin ovomucoid protease inhibitor solution was added to the single cell suspension, centrifuged at 300x g for 5min. Supernatant was removed, pellet resuspend in PBS and filtered through a 40um cell strainer (Pluristrainer; SKU 43-10040-50).

Filtered single cell suspension was centrifuged at 300x g for 5min at 4°C and resuspend in 100µl staining buffer (2%BSA/0.01%Tween, PBS) with 10µl Fc Blocking reagent (FcX, BioLegend) for 10 minutes on ice. 0.5µg of unique Cell Hashing antibodies were added and incubated for 20 minutes on ice. Stained cells were washed 3 times with 1mL staining buffer, spinned at 4 °C for 5 minutes at 350g. Stained single cell suspensions were counted for cell concentration and cell viability (TC20, Bio-rad). Typically, all single cell suspensions had a 70% or higher cell viability. Stained cells were pooled and re-counted until desired concentration for single cell sequencing (cell viability of at least 80%). Every sequencing round contained 14 hashed retinal organoid samples; the goal was to capture a total of approximately 30.000 cells per pool.

#### *Droplet-based single-cell RNA sequencing*

ScRNA-seq data was generated using the Chromium 10x 3'UTR-sequencing. Single cell suspensions were loaded onto the Chromium Single Cell system using the v3 chemistry. Subsequent steps were performed according to manufacturer's instructions.

#### *Computational analysis of single cell data*

Raw sequencing output were processed using the Cell Ranger (v6.0.1) pipeline (10X Genomics) with default settings and the pre-built human genome reference (GRCh38). Custom references of codon optimized AAV.h*CRB1*, codon optimized AAV.h*CRB2*, AAV.*GFP* and *CRB1-B* were added using the known FASTA sequence.

Filtered expression matrices were further processed with a Seurat (v4.1.0) based workflow in R (v4.1.0) (Hao et al., 2021). In short, cells were demultiplexed based on their HTO enrichment using HTODemux function of Seurat, and singlets were selected for downstream analysis. Quality control

followed, keeping cells with  $n\text{Feature\_RNA} > 800$  and  $< 6000$ ,  $n\text{Count\_RNA} < 30000$ , and  $\text{percent.mt} < 12$ . The raw counts were normalized with `NormalizedData` function (`scale.factor = 30000`). The top 2000 most variable genes were selected using `FindVariableFeatures`. Principal component analysis (PCA) was then performed using these 2000 genes. The first 15 PCs were used to calculate cell clusters and project the cells on a two-dimensional plot using Uniform Manifold Approximation and Projection (UMAP) algorithm. Top markers from `FindAllMarkers` function were analysed and compared to well-known cell type-specific markers to classify the clusters. For downstream analysis, data was subset per cluster, per patient derived retinal organoids, and/or per treatment. Then, differentially expressed genes were retrieved from the `FindMarkers` function. Genes with  $p\_val\_adj \leq 0,05$  were used for GO (Gene Ontology) term analysis, GO analysis was performed using `g:Profiler` (version `e106_eg53_p16_65fcd97`) with `g:SCS` multiple testing correction method applying significance threshold of 0.05 (Raudvere et al., 2019). Relevant terms and associated genes were included for visualization in this manuscript. For the violin plots comparing genes in *CRB1* patient-derived and isogenic control retinal organoids, the function `stat_compare_means` with a Wilcoxon t-test was used to determine statistically significant differences.

Number of cells used for downstream analysis per condition: (1) coming from one sequencing round, *ISO-P116-GFP*:  $n=993$  cells from 4 organoids, *P116-GFP*:  $n=1497$  cells from 4 organoids, *P116-CRB1-GFP*:  $n=1225$  cells from 3 organoids, *P116-CRB2-GFP*:  $n=884$  cells from 3 organoids, and (2) equally divided in three separate sequencing rounds, *ISO-P128-GFP*:  $n=5786$  cells from 6 organoids, *P128-GFP*:  $n=4908$  cells from 6 organoids, *P128-CRB1-GFP*:  $n=4386$  cells from 5 organoids, *P128-CRB2-GFP*:  $n=3870$  cells from 5 organoids, *P117-GFP*:  $n=3552$  cells from 5 organoids, *P117-CRB1-GFP*:  $n=3108$  cells from 5 organoids, *P117-CRB2-GFP*:  $n=2868$  cells from 5 organoids.

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