Human iPSC-Derived Retinas Recapitulate the Fetal CRB1 CRB2 Complex Formation and Demonstrate that Photoreceptors and Müller Glia Are Targets of AAV5

Peter M. Quinn,1,6 Thilo M. Buck,1,6 Aat A. Mulder,2 Charlotte Ohonin,1 C. Henrique Alves,1 Rogier M. Vos,3 Monika Bialecka,4 Tessa van Herwaarden,2 Elon H.C. van Dijk,1 Mays Talib,1 Christian Freund,1 Harald M.M. Mikkers,2 Rob C. Hoeben,2 Marie-José Goumans,2 Camiel J.F. Boon,1,5 Abraham J. Koster,2 Susana M. Chouva de Sousa Lopes,1 Carolina R. Jost,2 and Jan Wijnholds1,3,*

1Department of Ophthalmology, Leiden University Medical Center (LUMC), 2333 ZA Leiden, The Netherlands
2Department of Cell & Chemical Biology, Leiden University Medical Center (LUMC), 2333 ZA Leiden, The Netherlands
3Netherlands Institute for Neuroscience, Royal Netherlands Academy of Arts and Sciences (KNAW), 1105 BA Amsterdam, The Netherlands
4Department of Anatomy and Embryology, Leiden University Medical Center (LUMC), 2333 ZA Leiden, The Netherlands
5Department of Ophthalmology, Amsterdam University Medical Centers, Academic Medical Center, University of Amsterdam, 1000 AE Amsterdam, The Netherlands
6Co-first author
*Correspondence: j.wijnholds@lumc.nl
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SUMMARY

Human retinal organoids from induced pluripotent stem cells (hiPSCs) can be used to confirm the localization of proteins in retinal cell types and to test transduction and expression patterns of gene therapy vectors. Here, we compared the onset of CRB protein expression in human fetal retina with human iPSC-derived retinal organoids. We show that CRB2 protein precedes the expression of CRB1 in the developing human retina. Our data suggest the presence of CRB1 and CRB2 in human photoreceptors and Müller glial cells. Thus the fetal CRB complex formation is replicated in hiPSC-derived retina. CRB1 patient iPSC retinal organoids showed disruptions at the outer limiting membrane as found in Crb1 mutant mice. Furthermore, AAV serotype 5 (AAV5) is potent in infecting human Müller glial cells and photoreceptors in hiPSC-derived retinas and retinal explants. Our data suggest that human photoreceptors can be efficiently transduced by AAVs in the presence of photoreceptor segments.

INTRODUCTION

Mutations in the Crumbs homolog-1 (CRB1) gene are linked to an array of retinal dystrophies that exhibit high phenotypic variability and affect approximately 80,000 patients worldwide with an estimated prevalence in the US of 1/86,500 (Alves et al., 2014a; Stone et al., 2017; Talib et al., 2017). A meta-analysis has found that mutations in the CRB1 gene account for 2.7% and 10.1% of autosomal recessive retinitis pigmentosa (RP) and Leber congenital amaurosis (LCA) cases, respectively (Bujakowska et al., 2012). LCA is an early-onset disease, with newborns being blind around birth. However, we are yet to understand the localization of the CRB complex in early human fetal retinal development. No treatment is currently available for CRB1-associated retinal dystrophy in patients, but proof-of-concept gene supplementation studies have shown both morphological and functional rescue in CRB1 RP mouse models (Pellissier et al., 2015). A retrospective cohort study of patients with CRB1-associated RP has shown that gene therapeutic intervention is most likely required within the first three decades of life, but clinical endpoint criteria for a clinical trial need to be established from natural history studies (Talib et al., 2017). Preclinical considerations to be evaluated include the choice of adeno-associated virus (AAV) serotype for delivery of the clinical vector, in terms of potency, tropism, safety, and biodistribution.

Some retinal gene therapies use a serotype of AAV as a delivery vector in clinical trials (MacLaren et al., 2014; Maguire et al., 2008; Reichel et al., 2018) because it restricts vector tropism to specific retinal subpopulations, improves the efficiency of gene delivery, has low immunogenicity and long transgene expression. Cross-species differences in vector tissue tropism between mice and non-human primates have been previously highlighted and must be sufficiently addressed before moving toward clinical trials (Asokan et al., 2012; Ramachandran et al., 2017). We previously showed proof-of-concept for CRB1 gene therapy in Crb1-retinitis pigmentosa-like mice by subretinal application of an AAV9-CMV-CRB2 gene therapy vector, thereby demonstrating the need for transgene expression in both photoreceptors and Müller glial cells (MGCs) (Pellissier et al., 2015). On subretinal application, AAV9 and the AAV6 variant serotype ShH10Y445F are able to efficiently infect mouse photoreceptors, MGCs, and retinal pigment epithelium, whereas AAV5 does not efficiently infect and express in mouse MGCs (Aartsen et al., 2010; Pellissier et al., 2014a). AAV tropism differs between species, therefore the AAV serotype for clinical
gene therapy in both photoreceptors and MGCs needs to be validated. AAV5 and AAV9 infect non-human primate rod and cone photoreceptors (Boye et al., 2012; Vandenberghe et al., 2013). Human induced pluripotent stem cell (hiPSC)-derived retinal organoids, although in vitro, are a promising alternative or additional/pre-screening tool to animal models for evaluating transgene expression and biological activity (Quinn et al., 2018a). As previously demonstrated by others and ourselves hiPSC-derived retinal organoids and photoreceptors are amenable for the testing of AAV serotype/promoter combinations (Khabou et al., 2018; Quinn et al., 2018a; Wiley et al., 2016). However, for this purpose the hiPSC-derived retinal organoids should suitably recapitulate the human retina and the onset of expression of its proteins.

The core Crumbs complex in mammals is comprised of CRB1-3, PALS1 (also called MPP5), MUPP1, and PATJ. The prototypic CRB protein has a large extracellular domain with epidermal growth factor-like domains and laminin-A globular domains adjacent to a single transmembrane domain. A short C terminus of 37 amino acids contains an FERM protein-binding domain juxtaposing the single transmembrane domain. At the C-terminal end, there is a PDZ protein-binding motif of four amino acids (ERLI) that allows interaction with adaptor proteins such as PALS1 and PAR6 (Bachmann et al., 2001; Bulgakova and Knust, 2009; Lemmers et al., 2004; Roh et al., 2002). In non-human primates, CRB1 and CRB2 proteins localize to the subapical region adjacent to adherens junctions in MGCs and photoreceptors (Quinn et al., 2019). Similarly, in 2-day-old human adult cadaveric retina, both CRB1 and CRB2 are located at the subapical region in MGCs. However, in photoreceptors, the CRB1 protein is detectable at the subapical region near the outer limiting membrane (OLM), but CRB2 is not. In addition, both CRB1 and CRB2 are detected at vesicles in the photoreceptor inner segments at a distance from the OLM (Pellissier et al., 2014b, 2015). Thus far, the localization of the CRB complex in early human fetal retinal development is unknown.

In this study, we show the recapitulation of the CRB complex between the human fetal retina and cultured hiPSC-derived human retinal organoids. These studies highlight that CRB2, but not CRB1, is present at the subapical region in human fetal retina during the first trimester of pregnancy. CRB1 is expressed only at later time points from the second trimester onwards, concurring with the birth of differentiated cell types such as photoreceptors and MGCs. These data suggest role(s) for CRB2 but not CRB1 in the first trimester in the earliest human retinal radial glial progenitor cells. The data also suggest that CRB1 patient iPSC-derived retinal organoids develop a retinal phenotype as found in Crb1KO and Crb1KO/C249W mice (van de Pavert et al., 2004, 2007a, 2007b).

In addition, we show higher efficacy of AAV5 and ShH10Y445F over AAV9 serotypes for infection of photoreceptors in cultured human donor retinal explants. We also show the preference of AAV5 and ShH10Y445F over AAV9 to infect MGCs in hiPSC-derived retinal organoids. Overall, our results suggest that the AAV5 serotype, combined with CMV promoter-mediated expression, is suitable for gene therapy for CRB genes into human MGCs and rod and cone photoreceptors.
**RESULTS**

**Retinal Architecture in the Human Fetal Retina and iPSC Retinal Organoids**

The human fetal retina, as it transitions from the first to the second trimester of pregnancy, gives rise to all adult retinal cell types as the retina moves from a mitotic to post-mitotic state (Provis et al., 1985). We examined first (weeks 11–13) and second trimester (weeks 16–18) fetal retina and compared these with early (differentiation day 30 [DD30]) and late (DD120–DD240) healthy hiPSC retinal organoids. The retinal architecture in the human fetal retina recapitulated the retinal architecture as found in hiPSC retinal organoids (Figures S1 and S2).

**The CRB Complex in the Human Fetal Retina and iPSC Retinal Organoids**

We undertook immunohistochemistry studies to delineate the onset of expression of CRB1 and CRB2 in first and second trimester human fetal retina and in early and late stage differentiated retinal organoids.

In week 9 human fetal retina we did not detect the typical puncta-like CRB1+ immunostaining at the subapical region adjacent to the adherence junction marker β-catenin (Figures S3A–S3C). However, at week 11 we observed a gradient of CRB1 immunostaining located at the OLM (Figure 1A). In weeks 9 and 11, the human fetal retina stained positive for CRB2 at the subapical region adjacent to the adherence junctions as marked by anti-p120-catenin (Figures 1B and S3D–S3F). Furthermore, in the first trimester fetal retina at week 9 (Figures S3G–S3I), we found PATJ at the OLM and additionally in a subset of anti-Ki67+ cells; this was also seen in the developing mouse retina (Alves et al., 2013). In week 19, the human fetal retina expressed CRB1 (Figure 1E) and CRB2 (Figure 1F) with their prototypic puncta immunostaining pattern. Immunostaining for CRB complex members PALS1, MUPP1, PAR3, and the adherens junction markers β-catenin, p120-catenin, and N-cadherin were detected in both first and second trimester human fetal retina (Figure 1).

Similarly, in DD28 and DD80 retinal organoids we did not detect typical CRB1+ puncta-like immunostaining (Figures 1I and S3J). Its family member CRB2 was present at DD28 and DD80 (Figures 1J and S3K). PALS1 and MUPP1 were present at the subapical region in DD28 junctions being located in photoreceptor inner segments and apical villi of Müller glial cells. At least two independent samples were analyzed per time point. CC, connecting cilium; AJ, adherens junction; μ, microvilli; IS, inner segment. Scale bar (A–H), 1 μm; insert (D), 500 nm.
Moreover, in retinal organoids at DD30 (Figures S3N–S3P) we found PATJ at the OLM, and additionally in a subset of anti-Ki67+ cells. However, a typical and clear punctate-like staining pattern for CRB1 was detected at DD160 subapical of adherens junctions marker β-catenin (Figure 1K). CRB complex members CRB2, PALS1, MUPP1, and PAR3, and adherens junction markers p120-catenin and N-cadherin, were also detected in DD160 retinal organoids (Figures 1L–1O). CRB1 and CRB2 localization in retinal organoids was also confirmed in two other hiPSC lines: LUMC0080iCTRL12 (Figures S3Q–S3S) and LUMC0044iCTRL44 (Figures 5E–5H and 6E–6H).

**Ultra-Localization of CRB1 and CRB2 in the Human Fetal Retina and in iPSC Retinal Organoids**

We performed immunoelectron microscopy (immuno-EM) studies to analyze, at ultra-high resolution, the localization of CRB1 and CRB2 in photoreceptor cells (PRCs) and MGCs in first and second trimester human fetal retina and in retinal organoids. Immuno-EM for CRB1 in the first trimester fetal retina showed occasional and limited staining at putative inner segments (Figure 2A) and apical villi (Figure 2B) of radial glial progenitor cells. However, in the second trimester CRB1 labeling could be clearly detected at the subapical region adjacent to adherens junctions between putative photoreceptor inner segments (Figure 2C) and in the apical villi of radial glial progenitor cells/MGCs (Figure 2D). Immuno-EM for CRB2 showed pronounced labeling in first (Figures 3E and 3F) and second trimester (Figures 2G and 2H) fetal retina. CRB2 labeling localized at the plasma membrane and at the subapical region adjacent to adherens junctions between putative photoreceptor inner segments (Figures 2E and 2G) and in the apical villi of radial glial progenitor cells/MGCs (Figures 2F and 2H).

Similarly to first trimester human fetal retina, immuno-EM performed on retinal organoids showed sporadic and limited CRB1 labeling at the subapical region adjacent to adherens junctions between putative photoreceptor inner segments (Figure 3A) and in the apical villi of radial glial progenitor cells/MGCs (Figure 3B) in early retinal organoids. However, in late retinal organoids CRB1 labeling localized at the plasma membrane and at the subapical region adjacent to adherens junctions between putative photoreceptor inner segments (Figure 3C) and in the apical villi of radial glial progenitor cells/MGCs (Figure 3D). Immuno-EM for CRB2 showed pronounced labeling in early (Figures

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**Figure 3. CRB1 and CRB2 are Located at the Outer Limiting Membrane in Müller Glial Cells and Photoreceptors of Human iPSC-Derived Retinal Organoids**

Immuno-EM staining showing the localization of CRB1 (A–D) and CRB2 (E–H) in DD120 (A, B, E, and F) and DD170 (C, D, G, and H) LUMC0044iCTRL10 human iPSC-derived retinal organoids. CRB1 was lowly and sporadically detected at DD120 but being found subapically of adherens junctions adjacent to photoreceptor inner segments (A) (arrow) and in Müller glial cell apical villi (B) (arrowheads). At DD170, CRB1 was found consistently throughout the outer limiting membrane of human iPSC-derived retinal organoids subapically of adherens junctions of photoreceptors and Müller glial cells (C and D). CRB2 was localized at both DD120 and DD170 subapically of adherens junctions being located in photoreceptor inner segments and apical villi of Müller glial cells. At least two independent samples were analyzed per time point. CC, connecting cilium; AJ, adherens junction; μ, microvilli; IS, inner segment. Scale bar (A–H), 1 μm; inset (B and C), 500 nm.
Figure 3E and 3F) and late (Figures 3G and 3H) retinal organoids.

CRB2 labeling localized at the plasma membrane and at the subapical region adjacent to adherens junctions between putative photoreceptor inner segments (Figures 3E and 3G) and in the apical villi of radial glial progenitor cells/MGCs (Figures 3F and 3H).

Figure 4. CRB1 Patient Retinal Organoids Develop Proper Lamination

Immunohistochemistry pictures of healthy (LUMC0004CTRL10 and LUMC0044CTRL44) versus CRB1 patient (LUMC0116iCRB09, LUMC0117iCRB01, and LUMC0128iCRB01) retinal organoids at DD180. Sections were stained for pH3\(^+\) mitotic cells and Ki67\(^+\) cycling cells (A, E, I, M, and Q); Tuj1\(^+\) dendrites marking the GCL (B, F, J, N, and R); SOX9\(^+\) NBL cells (C, G, K, O, and S); and recoverin\(^+\) PRCs marking the ONL (D, H, L, P, and T). Experiments were validated in two differentiations for retinal organoids. NBL, neurobasal layer; GCL, ganglion cell layer. Scale bars (A–T), 20 µm. See also Figure S4.
Figure 5. CRB1 Patient Organoids Develop Retinal Degeneration
Immunohistochemistry pictures of healthy: LUMC0004iCTRL10 (A–D and A’–D’), and LUMC0044iCTRL44 (E–H, E’–H’), retinal organoids at DD180; and CRB1 patient: LUMC0116iCRB09 (I–L, I’–L’), LUMC0117iCRB01 (M–P, M’–P’, and W), LUMC0128iCRB01 (Q–T, Q’–T, (legend continued on next page)
Taken together, these data suggest that CRB1 is not required for the localization of CRB2, PALS1, MUPP1, or PATJ in the first trimester of human retinal development. The onset of CRB protein expression in human fetal retina is recapitulated in retinal organoids.

**Disruptions at the OLM in Retinal Organoids from CRB1 RP Patients**

We generated three hiPSC lines (LUMC0116iCRB, LUMC0117iCRB, and LUMC0128iCRB) from CRB1 RP patients. As with healthy iPSCs (Figure S4A), patient iPSCs were validated via immunostaining with pluripotent and germ layer markers (Figure S4B). LUMC0116iCRB has c.3122T > C p.(Met1041Thr) homozygote missense mutations. LUMC0117iCRB has 2,983G > T p.(Glu995*) and c.1892A > G, p.(Tyr631Cys) mutations. LUMC0128iCRB has c.2843G > A p.(Cys948Tyr) and c.3122T > C p.(Met1041Thr) missense mutations. The CRB1 gene mutations were re-confirmed in the established iPSCs (Figure S4C).

CRB1 patient iPSCs were able to differentiate to DD180 retinal organoids and were compared with healthy retinal organoids (Figure 4). The patient retinal organoids had pH3+ mitotic cells apically at the OLM, with Ki67+ cycling cells less tightly restricted to the middle of the neurobasal layer (NBL) (Figures 4I, 4M, and 4Q). The CRB1 patient iPSCs retinal organoids developed all three retinal layers: a ganglion cell layer marked by TuJ1+ dendrites (Figures 4J, 4N, and 4R), an NBL marked by SOX9+ retinal progenitor cells/MGCs (Figures 4K, 4O, and 4S), and an outer nuclear layer (ONL) marked by recoverin+ PRCs (Figures 4L, 4P, and 4T). However, frequent ectopic cells were found above the OLM (Figures 4I–4T). We observed many areas of funnel-shaped outward protruding recoverin+ PRCs (Figures 4L, 4P, and 4T) and, sporadically, SOX9+ cells above the OLM (Figure 4O).

All retinal organoids from the three different CRB1 patient lines developed small but frequent disruptions of CRB complex members at the OLM that were not detected in control lines (Figure 5). The adherens junction proteins N-cadherin, p120-catenin, and the subapical region proteins CRB2, PALS1, PAR3, and MUPP1 were localized as in healthy retinal organoids at DD180 (Figure 5). Interestingly, CRB1 variant protein localized similar to the wild-type CRB1 protein at the subapical region above the adherens junctions, but showed a curved and broadened expression pattern (Figures 5I, 5M, 5Q, and 5U) compared with the healthy control lines (Figures 5A and 5E). CRB1 variant protein was also mislocalized in the apical area of the NBL and in the ONL (Figures 5I, 5M, and 5Q). The mislocalized PRCs above the OLM resided at areas of OLM disruptions (Figures 5V, 5W, and 5X). In conclusion, the data from CRB1 patient hiPSC retinal organoids suggest a retinal degeneration phenotype similar to that previously found in mice lacking CRB1 or expressing variant CRB1C249W (van de Pavert et al., 2004, 2007b; 2007a).

**Transduction of Human iPSC Retinal Organoids with AAV5, AAV9, and ShH10Y445F**

Human iPSC-derived retinal organoids are a promising tool for evaluating transgene expression and biological activity (Quinn et al., 2018a). We have shown the need in Crb1-retinitis pigmentosa-like mice to direct CRB gene therapy to both photoreceptors and MGCs (Pellissier et al., 2015). Choosing the optimal promoter and AAV serotype for the therapeutic vector is therefore crucial to achieving expression in photoreceptors and MGCs. Here, we transduced the hiPSC-derived retinal organoids with AAV9-CMV-GFP, AAV5-CMV-GFP, ShH10Y445F-CMV-GFP, and ShH10Y445F-RLBP1-GFP at 10^{10} genome copies (gc). The CMV promoter drives expression of GFP in multiple cell types, whereas the hRLBP1 promoter drives expression in MGCs and retinal pigment epithelial cells (Pellissier et al., 2014a). When analyzed at the same laser intensity settings (Figures S5A–S5D) AAV5-CMV-GFP, ShH10Y445F-CMV-GFP, and ShH10Y445F-hRLBP1-GFP significantly outperformed AAV9-CMV-GFP at transducing DD220 hiPSC-derived retinal organoids collected 14 days after infection (Figures 6E and 6F). We quantified both the number of GFP+ cells per total cells (Figure 6E) and the mean gray value (Figure 6F) for each vector. The GFP+ nuclei were mainly located in the inner retina and exhibited radial projections. The GFP+ nuclei co-localized with anti-LHX2 (Figures 6A–6D, inserts) and anti-SOX2 (Figures S5E–S5H, and S5E–S5H), both transcription factors required for MGC development. Further proof of Müller glial-specific transduction is also seen with the use of ShH10Y445F-RLBP1-GFP (Figure 6D), which drives GFP in MGCs in rat and mouse retinas (Klimczak et al., 2009; Pellissier et al., 2014a). We additionally stained with anti-recoverin and found occasional co-localization between the photoreceptor marker and GFP+ nuclei (Figures S5I–SSSL and SS1′–SSS1′).

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Figure 6. AAV5 and SH10Y445F Efficiently Infect Müller Glial Cells in Human iPSC Retinal Organoids and Photoreceptors and Müller glial Cells in the Human Adult Retina

(A–F) Infection of 7.5-month-old LUMC0004iCTRL10 hiPSC retinal organoids at $10^{10}$ gc with AAV9-CMV-GFP (A), AAV5-CMV-GFP (B), SH10Y445F-CMV-GFP (C), and SH10Y445F-RLBP1-GFP (D), and co-staining with the Müller glial cell marker LHX2, the majority of LHX2+ cells were infected by AAV5 and SH10Y445F-RLBP1, whereas AAV9 and SH10Y445F-CMV-GFP had lower infection rates. (E) Bar graphs showing the percentage of GFP+ cells in total cells. (F) Fluorescence intensity (MGO) of infected cells. (G–N) Images of retina sections from different donors stained with ONL, INL, GCL, SOX9, RECOVERIN, and LHX2. (O–Q) Graphs showing the percentage of GFP+ cells in total cells and their localization in different layers of the retina.
In conclusion, these results indicate that AAV5 and ShH10Y445F serotypes are more potent transducers of MGCs than AAV9 in hiPSC-derived retinal organoids.

**Transduction of Adult Postmortem Human Retinal Explants with Serotypes AAV5, AAV9, and ShH10Y445F**

To verify the results of our transduction studies on hiPSC-derived retinal organoids we also tested AAV9-, AAV5-, and ShH10Y445F-CMV-GFP on human adult retinal explants. Initially, a titration study for AAV9 (Figures S6A–S6C), AAV5 (Figures S6D–S6F), and ShH10Y445F (Figures S6G–S6I) was undertaken to determine what gc level is required to infect photoreceptors and MGCs efficiently. Analysis for total infection (Figures S6P and S6Q), infection per retinal layer (Figures S6R and S6S), and tropism (Figure S6T) in donor 1 indicated 3 × 10^10 gc as a suitable level. Similar transduction patterns were found in donor 1 (Figures S6H–S6K, 6L–6N, S6B, S6D, and S6G), donor 2 (Figures S6J–S6L), and donor 3 (Figures S6M–S6O) at 3 × 10^10 gc. With individual analysis for total infection (Figures S6U, S6V, S6Y, and S6Z), infection per retinal layer (Figures S6W and S6AA), and tropism (Figures S6X and S6BB) also done for donors 2 and 3 at 3 × 10^10 gc. Donors 1–3 had similar retinal layer thickness (Figure S6CC) and cells per retinal layer (Figure S6DD). In the inner nuclear layer (INL), GFP co-labeled with MGC marker anti-SOX9 (Figures 6G–6J); and in the ONL, GFP co-labeled with photoreceptor marker anti-recoverin (Figures 6K–6N). When analyzing donors 1–3 together, AAV5 showed the higher efficacy of transducing retinal cell types than AAV9 (14% ± 5% versus 3% ± 1%; Figure 6O), and AAV5 and ShH10Y445F showed higher potency in transduction of photoreceptors in the ONL than AAV9 (11% ± 3% and 5% ± 1% versus 3% ± 1%; Figures 6P and 6Q). Interestingly, we noticed that the photoreceptors of cadaveric human retinal explants were only efficiently infected by AAV9 (Figures S7A and S7B), AAV5 (Figure S7C), or ShH10Y445F (Figure S7D) in the presence of intact photoreceptor segments (Figures S7E–S7G). This suggests an important role for the segments in the photoreceptor uptake of AAV particles. In conclusion, transduction of AAV serotypes in human cadaveric retina was more successful at targeting both photoreceptors and MGCs than in hiPSC-derived retinal organoids. AAV5 at 3 × 10^10 gc significantly outperformed AAV9 in the transduction of PRCs.

**CRB2 Is Located in the Apical Membrane of iPSC-Derived and Retinal Pigment Epithelium**

CRB2 but not CRB1 immunostaining was detected in first trimester human fetal retinal pigment epithelium (RPE) (Figures S3D’–S3F’). This was confirmed by immuno-EM in first and second trimester human fetal RPE (Figures 7A and 7B). CRB2 labeling was located above the adherens junctions at and above the tight junctions in the apical membrane and microvilli of human fetal RPE. Spheroids of hiPSC-derived RPE also generated during the differentiation method used (Liu et al., 2018; Quinn et al., 2018a; Zhong et al., 2014). These RPE spheroids initially attach to the periphery of the retinal organoids but can detach during culturing (Figures 7C, 7D, and 7D’). CRB1 could not be detected apically of β-catenin (Figure 7C) in hiPSC-derived RPE, but CRB2 was found apically of the adherens junction marker p120-catenin in DD160 hiPSC-derived RPE (Figure 7D). This pattern of localization was also found in hiPSC-derived RPE derived from hiPSC lines LUMC0080iCTRL12 (Figures 7E and 7F) and LUMC0044iCTRL44 (Figures 7G and 7H). Immuno-EM of iPSC-derived RPE confirmed the apical staining for CRB2 above adherens junctions at and above the tight junctions in the apical membrane and microvilli. Aspecific staining was detected within melanin granules due to the presence of endogenous peroxidase in these structures (Figures 7I and 7I’). Electron microscopy of hiPSC-derived RPE also

**Scale bar (A–D, G–N), 20 μm; (A–D, G–N) inserts, 10 μm. Data are presented as means ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001. See also Figures S5–S7.**
showed the presence of melanosomes with pigments, basally located mitochondria and basement membrane (Figure 7J, insert), and fibrous long-spacing collagen (Figures 7J and 7J'). RPE cells were also infected by AAV9, AAV5, and ShH10Y445F (Figures S7H–S7K).

DISCUSSION

In this study, we showed (1) that in human fetal retina during the first trimester of pregnancy CRB2 is the predominant CRB family member in radial glial progenitor cells; and that CRB1 onset of expression at the subapical region coincides with the maturation of the retina during the second trimester. (2) CRB2 but not CRB1 is expressed in the fetal RPE. (3) The onset of CRB protein expression in human fetal retina and RPE is recapitulated in hiPSC-derived retinal organoids and RPE. (4) CRB1 RP patient retinal organoids develop disruptions at the OLM with misplaced photoreceptors. (5) AAV5 and ShH10Y445F serotypes are more potent than AAV9 serotype in infecting cultured retinal organoids. (6) AAV5-CMV-GFP is more efficient than AAV9-CMV-GFP to express GFP in PRCs in cultured human donor retinal explants. (7) Human PRCs are efficiently transduced only in the presence of photoreceptor segments.

In the human fetal retina, we found that CRB2 is the predominant CRB effector protein in the first trimester of pregnancy. CRB2 is a gene expressed in several tissues, including the cerebral cortex (Dudok et al., 2016), with a crucial role during early development in both mice and humans. Mice lacking Crb2 are embryonic lethal, with a critical role for the CRB2 protein during gastrulation in the epithelial-to-mesenchymal transition (Ramkumar et al., 2016). CRB2 protein variants in humans have been linked to a syndromic phenotype causing kidney and brain dysfunctions and lethality (Ebarasi et al., 2015; Slavotinek et al., 2015) as well as to RP (Chen et al., 2018). In the second trimester, CRB1 and CRB2 localized at the subapical region in apical villi of radial glial progenitor cells/MGCs and at the subapical region above the adherens junctions in the inner segments of PRCs. During the second trimester, the

$\text{CRB1}$ RP patient retinal organoids develop disruptions at the OLM with misplaced photoreceptors. (5) AAV5 and ShH10Y445F serotypes are more potent than AAV9 serotype in infecting cultured retinal organoids. (6) AAV5-CMV-GFP is more efficient than AAV9-CMV-GFP to express GFP in PRCs in cultured human donor retinal explants. (7) Human PRCs are efficiently transduced only in the presence of photoreceptor segments.
retina undergoes the birth of all adult cell types, and the retina is transitioning from a mitotic to post-mitotic state (Lee et al., 2006; Provis et al., 1985). Retinal organoids go from an early highly cell-cycling state, in which Ki67 marks the entire NBL at DD28, toward a moderate cell-cycling state, in which Ki67 becoming restricted to the mid-NBL at DD120.

Interestingly, here we showed that the onset of CRB1 protein expression coincided with the maturation of the retinal organoids, and this finding is recapitulated in the human fetal retina. In early-stage retinal organoids, we found, as in the first trimester fetal retina, CRB2 but little CRB1 protein expression at the subapical region. In later-stage hiPSC-derived retinal organoids we found CRB2 and CRB1 protein expression at the subapical region as in second trimester fetal retina. We also found a recapitulation of CRB2 expression when comparing first trimester fetal RPE with hiPSC-derived RPE.

We present here the generation and characterization of CRB1 patient-derived hiPSCs differentiated to retinal organoids. We demonstrate that patient retinal organoids give rise to a morphological significant phenotype even though variant CRB1 protein and its interaction partners (MUPP1, PALS1, and CRB2) are detected at the OLM. The data suggest disruptions at the OLM resulting in loss of adhesion between photoreceptors and MGCs. Decreased levels of CRB1 and CRB2 proteins at the OLM exacerbated retinal degeneration in mouse models (Alves et al., 2013; Pellissier et al., 2013; Quinn et al., 2018b). Also, the volcanic-like cell protrusions and OLM disruptions in the patient retinal organoids show striking similarities to the morphological phenotype found in 3-month-old Crb1KO and 8-month-old Crb1KO/C249W RP mice (van de Pavert et al., 2004, 2007b; 2007a). Further studies are needed to elucidate the underlying effects of the variant CRB1 proteins on protein-protein interactions and downstream cell signaling pathways.

We hypothesize that retinal organoids could be a good model for evaluating transgene expression and biological activity due to their close mimicking of human fetal retinal development (Quinn et al., 2018a). Our transduction studies on cadaveric human retinal explants showed a higher potency for AAV5 over AAV9 for transduction of photoreceptors. Also, the data suggest the higher efficacy of AAV5-CMV-GFP than AAV9-CMV-GFP or ShH10Y445F-CMV-GFP to express in PRCs and MGCs. In the absence of photoreceptor segments in the human retinal explants, AAV5-CMV-GFP, ShH10Y445F-CMV-GFP, and AAV9-CMV-GFP showed higher efficacy to express in MGCs than in PRCs. The latter tropism and expression potency data in cultured cadaveric human retinal explants are reproduced in retinal organoids that recapitalize second trimester fetal retina.

Previous subretinal injection studies in which AAV5 was administered in mice at postnatal day 0 (P0) or P30 have shown preferred transduction of P0 cone PRC and MGCs, but only of P30 rod and cone PRCs (Surace et al., 2003). We hypothesize that this preference in transduction patterns of PRC and MGCs in immature versus mature retina is due to the presence or absence of matured photoreceptor segments. In mice, photoreceptor segments seem to be required for the efficient transduction of PRCs with AAV vectors (Petit et al., 2017). A very interesting and clinically relevant finding is that photoreceptors in cultured cadaveric human retinal explants are only efficiently transduced when they have photoreceptor segments. Retinal organoids represent immature fetal retinas that contain PRCs but with yet very immature segments. We hypothesize that PRCs are transduced by AAV5, AAV9, and ShH10Y445F once the PRC segments are formed in sufficient number and size. In the absence of PRC segments, however, there is increased bioavailability of AAV vectors to target less-abundant/pREFERRED receptors for AAV uptake, e.g., on MGCs. Interestingly, dependency on the presence of photoreceptor segments for photoreceptor transduction was observed for all three AAV serotypes (AAV5, AAV9, and ShH10Y445F), suggesting a putative common mechanism of active AAV uptake into photoreceptors. The inner segments are a putative site of receptor-dependent or –independent clathrin- and caveolae-mediated endocytosis (Fuchs et al., 2014).

Our data suggest that for clinical gene therapy with AAV5, AAV9, or ShH10Y445F the target PRCs should have intact photoreceptors to become efficiently transduced. It also implies that the AAV vector particles should be able to reach the PRC segments during clinical surgical application. This condition of accessibility of PRC inner segments in human retina in vivo is met upon subretinal injection, as suggested by AAV5 or AAV9 infection of PRCs in non-human-primate retinas (Boye et al., 2012; Vandenberghe et al., 2011). Our mice lacking CRB1, as well as the mice with reduced levels of CRB2, showed a compromised OLM. We further hypothesize from our previous studies in mice that the retinas of human patients with loss of CRB1, or expressing non-functional CRB1 variants, have a compromised OLM that allows increased passage of AAV viral particles across the adherence junctions to reach the AAV-receptor molecules on MGCs (Pearson et al., 2010).

**EXPERIMENTAL PROCEDURES**

See further details in the Supplemental Experimental Procedures.

**Fetal Human Retinal Tissue**

The use and collection of the material was approved by the Medical Ethics Committee of the Leiden University Medical Center (P08.087).
Adult Human Retinal Tissue
Tissue was collected in agreement with the guidelines of the ethics committee of the LUMC. Informed consent was obtained on the basis of the Declaration of Helsinki (World Medical Association).

Cell Culture and Retinal Organoid Differentiation
Human iPSCs (LUMC0004iCTRL10 [Dambrot et al., 2014], LUMC0044iCTRL44 [Chen et al., 2017], LUMC0080iCTRL12 [Figure S4A], LUMC0116iCRB09, LUMC0117iCRB01, and LUMC0128iCRB01 [Figure S4B]) were maintained on Matrigel (BD)-coated plates in mTeSR medium (STEMCELL Technologies) and passaged mechanically. Retinal organoid differentiation was carried out as previously reported (Quinn et al., 2018a; Zhong et al., 2014).

Electron Microscopy
Immuno-EM was performed as described previously (Klooster et al., 2011). In brief, sections were incubated with antibody for 48 h, then incubated with appropriate secondary peroxidase anti-peroxidase for 2 h, then developed in a 2,2-diaminobenzidine solution for 4 min, and then the gold substitute silver peroxidase method applied.

Generation and Purification of the Viral Vectors
The pAAV2-eGFP plasmids were generated previously and consist of the flanking inverted terminal repeats of AAV2, the full-length CMV promoter, or the human RLB1 promoter, the eGFP cDNA, the Woodchuck posttranscriptional regulatory element, and the bovine growth hormone poly(A) (Aartsen et al., 2010; Alves et al., 2014b; Pellissier et al., 2014a).

In Vitro Transduction of Human Donor Retina and Human Induced Pluripotent Stem Cell-Derived Retinal Organoids
In vitro transduction protocols for (1) human donor retina and (2) hiPSC-derived retinal organoids have been described previously (Buck et al., 2018; Quinn et al., 2018a).

Statistical Method
All statistical analyses were performed using GraphPad Prism version 7 (GraphPad Software). All values are expressed as mean ± SEM. Multiple t tests were performed with the analyzed number of samples indicated in the figure legends. Immunohistochemistry was performed on iPSC-derived retinal organoids from three independent healthy and three independent CRB1-patient iPSC lines from two or more differentiations, with three to six sections examined per organoid. Immunohistochemistry was performed on at least two independent human fetal eyes per time point, with three to six sections examined per eye. Immuno-EM was performed on at least two independent human fetal eyes, iPSC-derived retinal organoids, and RPE for each time point. For AAV transduction studies, between three and six different sections from at least three different human donor retina or hiPSC-derived retinal organoids (one to two organoids from two independent differentiations analyzed) were used for quantification. Between three and five images per organoid and ten images per adult donor retina were analyzed.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.stemcr.2019.03.002.

AUTHOR CONTRIBUTIONS
P.M.Q., T.M.B., and J.W. conceived and designed the experiments. P.M.Q., T.M.B., C.O., C.H.A., and A.A.M. performed the experiments. R.M.V. produced virus stocks. M.B., T.v.H., E.H.C.v.D., and M.T. collected study material. H.M.M.M., C.F., R.C.H., M.-J.G., C.J.F.B., A.J.K., and S.M.C.d.S.L. provided study material and/or access to facilities. P.M.Q. and T.M.B. assembled the data. P.M.Q., T.M.B., A.A.M., C.R.J., and J.W. analyzed and interpreted the data. P.M.Q., T.M.B., and J.W. wrote the manuscript. All authors reviewed the manuscript. J.W. provided funding acquisition, supervision, and final approval of manuscript.

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